# 08/876,276 Search Strategy/Results

(FILE 'HOME' ENTERED AT 08:22:03 ON 03 MAY 2002) FILE 'CAPLUS' ENTERED AT 08:22:11 ON 03 MAY 2002 E ZHANG/AU 25

FILE 'MEDLINE, AGRICOLA, CAPLUS, BIOSIS, EMBASE, WPIDS' ENTERED AT 08:29:22 ON 03 MAY 2002
149646 S FLOW (W) CYTOMETRY
866 S L1 AND LIBRARY
392 S L2 NOT PY-1997
107 S L3 AND (FLUORESCENCE OR FLUORESCENT)
66 DUP REM L4 (41 DUPLICATES REMOVED)

L1 L2 L3 L4 L5

FILE

ANSWER 1 OF 66 DUPLICATE 1 MEDLINE

ACCESSION NUMBER: MEDLINE 97405894 PubMed ID: 9260938 DOCUMENT NUMBER: 97405894

97405894 PubMed 1D: 9280938
The glycolytic enzyme glyceraldehyde-3-phosphate dehydrogenase of Candida albicans is a surface antigen.
Gil-Navarro I; Gil M L; Casanova M; O'Connor J E; Martinez J P; Gozalbo D
Departamento de Microbiologia y Ecologia, Facultad de

AUTHOR:

CORPORATE SOURCE:

SOURCE:

Departamento de Microbiologia y Ecologia, Facultad d Farmacia, Universitat de Valencia, Spain. JOURNAL OF BACTERIOLOGY, (1997 Aug) 179 (16) 4992-9. JOURNAL code: HH3; 2985120R. ISSN: 0021-9193. United States

PUB. COUNTRY:

Journal; Article; (JOURNAL ARTICLE)

LANGUAGE: English

Priority Journals GENBANK-U72203 FILE SEGMENT: OTHER SOURCE: ENTRY MONTH:

ENTRY DATE:

RR SOURCE: GENBANK-U72203
RY MONTH: 199709
RY DATE: Entered STN: 19970922
Last Updated on STN: 19970905
A lambda gtll cDNA library from Candida albicans ATCC 26555 was screened by using pooled sera from two patients with systemic candidiasis and five neutropenic patients with high levels of anti-C. albicans immunoglobulin M antibodies. Seven clones were isolated from 60,000 recombinant phages. The most reactive one contained a 0.9-kb cDNA encoding a polypeptide immunoreactive only with sera from patients with systemic candidiasis. The whole gene was isolated from a genomic library by using the cDNA as a probe. The nucleotide sequence of the coding region showed homology (78 to 79%) to the Saccharomyces cerevisiae TDM1 to TDH3 genes coding for glyceraldehyde-3-phosphate dehydrogenase (GAPDH), and their amino acid sequences showed 76% identity; thus, this gene has been named C. albicans TDH1. A rabbit polyclonal antiserum against the purified cytosolic C. albicans GAPDH (polyclonal antibody [PAD] anti-CA-GAPDH) was used to identify the GAPDH in the beta-mercaptoethanol extracts containing cell wall moieties. Indirect immunofluorescence demonstrated the presence of GAPDH at the C. albicans cell surface, particularly on the blastoconidia. Semiquantitative flow cytomatry analysis showed the sensitivity of this GAPDH form to trypsin and its resistance to be removed with 2 M NaCl or 2% sodium dodecyl sulfate. The decrease in fluorescence in the presence of soluble GAPDH indicates the specificity of the labelling. In addition, a dose-dependent GAPDH enzymatic activity was detected in intact blastoconidia and germ tube cells. This activity was detected in intact blastoconidia and germ tube cells. This activity was reduced by pretreatment of the cells with trypsin, formaldehyde, and PAD anti-CA-GAPDH. These observations indicate that an immunogenic, enzymatically active cell wall-associated form of the glycolytic enzyme GAPDH is found at the cell surface of C. albicans cells.

ANSWER 2 OF 66 MEDLINE DUPLICATE 2

1998173052 MEDLINE 98173052 PubMed ID: 9514119 ACCESSION NUMBER: DOCUMENT NUMBER:

TITLE:

981/3052 Pubmed ID: 9514119 Isolation of anti-T cell receptor scFv mutants by yeast surface display. Kieke M C; Cho B K; Boder E T; Kranz D M; Wittrup K D Department of Biochemistry, University of Illinois, Urbana 61801 AUTHOR:

CORPORATE SOURCE:

CONTRACT NUMBER:

ROLAIS990 (NIAID)
T32 GM07283 (NIGMS)
PROTEIN ENGINEERING, (1997 Nov) 10 (11) 1303-10.
Journal code: PR1; 8801484. ISSN: 0269-2139.
ENGLAND: United Kingdom
Journal; Article; (JOURNAL ARTICLE) SOURCE:

PUB. COUNTRY:

LANGUAGE: English

FILE SEGMENT: ENTRY MONTH:

Priority Journals 199804

ENTRY DATE:

INSMERN: Priority Journals
ISAGENT: Entered STN: 19980507
Entered Medline: 19980424
Yeast surface display and sorting by flow cytometry
have been used to isolate mutants of an scPv that is specific for the
Vbeta8 region of the T cell receptor. Selection was based on equilibrium
binding by two fluorescently labeled probes, a soluble Vbeta8 domain and
an antibody to the c-myc epitope tag present at the carboxy-terminus of
the scPv. The mutants that were selected in this screen included a scPv
with threefold increased affinity for the Vbeta8 and scFv clones that were
bound with reduced affinities by the anti-c-myc antibody. The latter
finding indicates that the yeast display system may be used to map
conformational epitopes, which cannot be revealed by standard peptide
screens. Equilibrium antigen binding constants were estimated within the
surface display format, allowing screening of isolated mutants without
necessitating subcloning and soluble expression. Only a relatively small
library of yeast cells (3 x 10[5]) displaying randomly mutagenized
scPv was screened to identify these mutants, indicating that this system
will provide a powerful tool for engineering the binding properties of
eucaryotic secreted and cell surface proteins.

ANSWER 3 OF 66 MEDLINE

ACCESSION NUMBER:

DOCUMENT NUMBER:

MEDLINE
97452346 MEDLINE
97452346 PubMed ID: 9308760
Identification of an epitope of alpha-enolase (a candidate plasminogen receptor) by phage display.
Arza B; Felez J; Lopez-Alemany R; Miles L A; Munoz-Canoves TITLE:

AUTHOR:

CORPORATE SOURCE: Institut de Recerca Oncologica, Hospital Duran i Reynals,

Barcelona, Spain. HL-38272 (NHLBI) HL-45934 (NHLBI) CONTRACT NUMBER:

THROMBOSIS AND HAEMOSTASIS, (1997 Sep) 78 (3) 1097-103. JOURNAL code: VQ7; 7608063. ISSN: 0340-6245. GERMANY: Germany, Federal Republic of Journal; Article; (JOURNAL ARTICLE) SOURCE:

PUB. COUNTRY:

LANGUAGE: English Priority Journals

FILE SEGMENT: ENTRY MONTH: 199801 ENTRY DATE: Entered STN: 19980129 Last Updated on STN: 19980129 Entered Medline: 19980114

Last Updated on STN: 19980129

Entered Medline: 19980114

Alpha-enolase is an ubiquitous cytoplasmic glycolytic enzyme which also exhibits cell surface mediated functions and a structural role in the lens of some species. An alpha-enolase related molecule (alpha-ERM) is present on the surfaces of neutrophils, monocytes and monocytoid cells and has the capacity to specifically bind plasminogen, suggesting that alpha-ERM may function as a plasminogen receptor. We have generated a monoclonal antibody (mAB), 9C12, against alpha-ERM. This mAB reacted with both alpha-ERM and purified human alpha-enolase in Western blotting and in enzyme linked immunosorbent assays (ELISA). mAB 9C12 detected a cell surface associated molecule on human peripheral blood neutrophils and on U937 human monocytoid cells as assessed by fluorescence activated cell sorting (FACS) analyses. In addition, mAB 9C12 recognized an intracellular pool of alpha-enolase/alpha-ERM in permeabilized U937 cells. A phage display approach was employed to identify the alpha-enolase epitope recognized by mAB 9C12. Random fragments of 100-300 base pairs (bp), obtained from the full length human alpha-enolase ChAM, were cloned into the filamentous phage vector pComb3B, to generate a phage-displayed peptide library. Recombinant phages binding to mAB 9C12 were selected and their DNA inserts characterized by direct sequencing. All of the fragments which bound to mAB 9C12 encoded the common sequence DLDFKSPDDPSRYISP, spanning amino acids 257-272 of human alpha-enolase. This sequence is located within an external loop of the molecule. These data indicate that this sequence contains the epitope recognized by mAB 9C12 and is, therefore, exposed on the cell surface, further suggesting that alpha-enolase and alpha-ERM share common amino acid sequences.

ANSWER 4 OF 66 MEDLINE

ACCESSION NUMBER: DOCUMENT NUMBER: 97325526 MEDLINE

97325526 PubMed ID: 9181579
Use of a modified bacteriophage to probe the interactions between peptides and ion channel receptors in mammalian TITLE:

cells. Li M

AUTHOR: CORPORATE SOURCE:

Department of Physiology, Johns Hopkins University School of Medicine, Baltimore, MD 21205, USA.. min-li@qmail.bs.jhu.edu
NATURE BIOTECHNOLOGY, (1997 Jun) 15 (6) 559-63.
JOURNAL COde: CQ3; 9604648. ISSN: 1087-0156.

SOURCE:

PUB. COUNTRY: United States

Journal; Article; (JOURNAL ARTICLE) LANGUAGE:

English FILE SEGMENT: ENTRY MONTH:

Priority Journals 199708 ENTRY DATE:

Y MONTH: 199708
Y DATE: Entered STN: 19970902
Last Updated on STN: 19970902
Entered Medline: 19970819
Besides natural peptide ligands, screening of random peptide libraries has yielded novel bloactive peptides for cell surface receptors. A method is described that uses a modified bacteriophage as a detection reagent to monitor the expression of receptor channels in mammalian cells and to probe the molecular interaction between phare-tethered peptides (shi monitor the expression of receptor channels in mammalian cells and to probe the molecular interaction between phage-tethered peptides (phi T-peptides) and specific receptor targets. By taking advantage of a specific multivalent interaction between phi T-peptides and the receptor target, assays have been developed that use phi T-peptides specific for the N-methyl-D-aspartate glutamate receptor, an important ligand-gated ion channel in the nervous system, to monitor the receptor expression in cultured mammalian cells. Combining these phi T-peptide binding assays with fluorescence-activated cell sorting, 10(4) random glutamate receptor mutants were screened and candidate interaction residues were identified. This dual heterologous expression system offers a powerful approach to the molecular studies of protein-protein interactions.

```
ANSWER 5 OF 66
                   MEDLINE
                                                    DUPLICATE 3
```

ACCESSION NUMBER: DOCUMENT NUMBER: 97222477 97222477 MEDLINE PubMed ID: 9069120

TITLE:

97222477 PubMed ID: 9069120
Mapping of microsatellite markers developed from a flow-sorted swine chromosome 6 library.
Grimm D R; Goldman T; Holley-Shanks R; Buoen L; Mendiola J; Schook L B; Louis C; Rohrer G A; Lunney J K
USDA-ARS-IDRL, BARC-East, Building 1040, Room 105, Beltsville, MD 20705, USA.
MAMMALIAN GENOME, (1997 Mar) 8 (3) 193-9.
Journal code: BES; 9100916. ISSN: 0938-8990.
United States AUTHOR:

CORPORATE SOURCE:

SOURCE:

PUB. COUNTRY: United States

Journal; Article; (JOURNAL ARTICLE)
English

LANGUAGE:

FILE SEGMENT: Priority Journals

OTHER SOURCE:

GENBANK-U24283; GENBANK-U24284; GENBANK-U24285; GENBANK-U24286; GENBANK-U62944; GENBANK-U62945; GENBANK-U62946; GENBANK-U62947; GENBANK-U62948; GENBANK-U62949; GENBANK-U62950; GENBANK-U62951;

GENBANK-U62952; GENBANK-U62953; GENBANK-U62954; GENBANK-U62955; GENBANK-U62956; GENBANK-U62957; GENBANK-U62958; GENBANK-U62959; GENBANK-U66700

ENTRY MONTH: 199706

ENTRY DATE:

GENBANK-U62958; GENBANK-U62959; GENBANK-U66700
Y MONTH: 199706
Y DATE: Entered STN: 19970630
Last Updated on STN: 19970619
Swine Chromosome (Chr) 6-enriched libraries, generated with size-fractionated DNA isolated from chromosomes sorted by flow cytometry, have been used to develop new Chr 6 microsatellite markers. Chromosome isolation procedures were established to reproducibly prepare high quality chromosomes from phytohemagglutinin (PHA)-stimulated swine peripheral blood lymphocytes and to sort individual chromosomes after staining with Hoechst 33258 and chromosycin A3. Chromosome purity was verified by specific staining of swine Chr 6 with fluorescence in situ hybridization (FISH) by use of painting probes generated by degenerate oligonucleotide-primed polymerase chain reaction (DOP-PCR) amplification of as few as 300 sorted Chr 6. For library construction, DNA was extracted from flow-sorted pools representing Chr 6, amplified, size selected for fragments from 300 to 700 bp, and ligated

into pBluescript SK II+ or Lambda ZAP Express. The libraries were then screened with a radiolabeled poly-(dCA) DNA probe. Of 107 (CA)n repeat-containing clones verified by sequencing, 21 were polymorphic and used to genotype the University of Illinois swine reference families. Linkage analysis was then performed with CRIMAP 2.4 (LOD > 3.0), and the results showed that 15 of the microsatellites mapped to swine Chr 6. At least three of these new markers map to locations where there were gaps in the consensus Chr 6 map. Another four markers, because of their PIC values, should provide more informative markers in other areas of the map. Most of the new markers can also be used for automated genotyping with fluorescent labeling. This set of 15 new Chr 6 markers will, therefore, be useful in helping to define QTL associated with swine Chr 6.

L5 ANSWER 6 OF 66 EMBASE COPYRIGHT 2002 ELSEVIER SCI. B.V. ACCESSION NUMBER: 97088951 EMBASE 1997088951 Cell display library for gene cloning of variable DOCUMENT NUMBER: Cell display library for gene cloning of variable regions of human antibodies to hepatitis B surface antigen. Higuchi K.; Araki T.; Matsuzaki O.; Sato A.; Kanno K.; Kitaguchi N.; Ito H. K. Kanno, Pharmaceutical Group, Fundamental Res. Lab. of Life Sci., Asahi Chemical Industry Co., Ltd., 2-1 Samejima, Fuji, Shizuoka 416, Japan. a8012181@ut.asahi-kasei.co.jp Journal of Immunological Methods, (1997) 202/2 (193-204). Refs: 27 CORPORATE SOURCE: SOURCE: ISSN: 0022-1759 CODEN: JIMMBG S 0022-1759(97)00010-0 PUBLISHER IDENT .: Netherlands
Journal; Article
004 Microbiology
022 Human Genetics DOCUMENT TYPE: FILE SEGMENT: 026 Immunology, Serology and Transplantation LANGUAGE English SUMMARY LANGUAGE: English A novel cell display system was developed for cloning the variable region (V) genes of antigen-specific human antibodies. The system is based on an (V) genes of antigen-specific human antibodies. The system is based on an antibody library displayed on the surface of COS cells, using a plasmid vector designed to direct expression of membrane-bound antibodies. COS cells expressing antigen-specific antibodies were separated using a flow cytometer for their binding to a fluorescent dye-labeled antigen. To test the performance of this system, we cloned V genes of 4 antibodies directed against hepatitis B surface antigen (HBsAg) from a library prepared from peripheral blood lymphocytes of a vaccinated donor. These membrane-bound anti-HBsAg antibodies were easily converted to soluble forms, all of which showed a size similar to human serum IgG in SDS-PAGE and the same specific binding to HBsAg as membrane-bound forms in ELISA. All V(H) and Y(.kappa.) gene segments of the 4 clones isolated in this study belonged to V(H)III and V(.kappa.) I subgroups, respectively. These findings demonstrate the potential and selection capabilities of our These findings demonstrate the potential and selection capabilities of our cell display system for cloning the V genes of antigen-specific human antibodies.

ANSWER 7 OF 66 MEDLINE DUPLICATE 4

97362024 ACCESSION NUMBER: MEDITINE DOCUMENT NUMBER:

PubMed ID: 9215560 TITLE:

97382024 PLOMENT ID: 9215560
Chromosomal homeologies between human, harbor seal (Phoca vitulina) and the putative ancestral carnivore karyotype revealed by Zoo-FISH.
Pronicke L; Muller-Navia J; Romanakis K; Scherthan H
Abteilung fur Humanbiologie und Humangenetik, Universitat

AUTHOR .

CORPORATE SOURCE: Kaiserslautern, Postfach 3049, D-67653 Kaiserslautern,

Germany. CHROMOSOMA, (1997 Jul) 106 (2) 108-13 SOURCE:

Journal code: D7A; 2985138R. ISSN: 0009-5915. GERMANY: Germany, Federal Republic of Journal; Article; (JOURNAL ARTICLE) PUB. COUNTRY:

LANGUAGE: FILE SEGMENT: ENTRY MONTH: Priority Journals 199709 Entered STN: 19970922

Y DATE: Entered STN: 19970922
Last Updated on STN: 19970992
Entered Medline: 19970909
We report on the construction of the first comparative Zoo-FISH map of a marine mammal. Zoo-FISH with DNA probes from a human chromosome-specific library to metaphase spreads of the harbor seal (Phoca vitulina) disclosed 31 conserved syntenic segments covering the complete autosomal complement and the X chromosome. Comparison with Zoo-FISH maps of other species reveals that the harbor seal shares a high degree of karyotypic homeology with the human complement and an even higher degree with the conordinal cat complement. These findings suggest that pinniped, felid and human karyotypes have maintained conserved complements. Based on data of Zoo-FISH and comparative cytogenetics, a Zoo-FISH map of the ancestral carnivore karyotype (Z-CAR) is proposed. Flow cytometry revealed that the DNA value of the harbor seal genome is 79% that of the human genome. human genome.

L5 ANSWER 8 OF 66 CAPLUS COPYRIGHT 2002 ACS ACCESSION NUMBER: 1997:333943 CAPLUS

1997:333943 CAPLUS 127:3859

DOCUMENT NUMBER:

An optimized method for cell-based phage display

AUTHOR (S):

Watters, Jennifer M.; Telleman, Pieter: Junghans.

Richard P.

Biotherapeutics Development Lab, Department of Medicine, Harvard Medical School, New England Deaconess Hospital, Boston, MA, 02215, USA Immunotechnology (1997), 3 (1), 21-29 CODEN: IOTEER; ISSN: 1380-2933 Elsevier
Journal CORPORATE SOURCE:

SOURCE:

PUBLISHER DOCUMENT TYPE: Journal LANGUAGE: English

Traditional methods of phage display panning, bind purified antigen to plates or other solid phases to which libraries are then applied, followed by vigorous washings in detergent-supplemented buffers to select for

specific phage Fab. These methods are not directly applicable to antigens in their native environment on cell surfaces or in settings where the target antigen is unknown. To develop a model antigen system employing target antigen is unknown. To develop a model antigen system employing whole cells rather than purified protein immobilized on a substrate; to optimize methods for phage display panning using a cell-based system. Specificity of binding of phage Fab to antigen on cells was demonstrated by output titer and by flow cytometry. Output titers showed a plateau and binding advantage, after four washes, corresponding to removal of most non-specifically bound phage. Enrichment advantage was independent of input phage no. Longer incubation times, to cell tolerance, improved specific binding. Temp. had modest impact as a variable in the panning and washing. An increase in output titers paralleled enrichment for specific phage Fab. An optimized method applied to whole cells can productively enrich specific phage Fab in mixts. with large excesses of non-specific phage Fab over several rounds of panning.

ANSWER 9 OF 66 MEDLINE

ACCESSION NUMBER: DOCUMENT NUMBER: 96279212 96279212 MEDLINE

96279212 MEDLINE
96279212 PubMed ID: 8663178
Rapid identification of phosphopeptide ligands for SH2
domains. Screening of peptide libraries by
fluorescence-activated bead sorting.
Muller K; Gombert F O; Manning U; Grossmuller F; Graff P;
Zaegel H; Zuber J F; Freuler F; Tschopp C; Baumann G
Sandoz Pharma Ltd., Preclinical Research, CH-4002 Basel,
Switzerland. AUTHOR:

CORPORATE SOURCE:

JOURNAL OF BIOLOGICAL CHEMISTRY, (1996 Jul 12) 271 (28) SOURCE:

16500-5. Journal code: HIV; 2985121R. ISSN: 0021-9258.

PUB. COUNTRY:

United States
Journal; Article; (JOURNAL ARTICLE)

LANGUAGE:

English Priority Journals 199608

FILE SEGMENT: ENTRY MONTH:

ENTRY DATE:

SEGMENT: Priority Journals

Y MONTH: 199608

Y MONTH: 199608

Thered STN: 19960911

Entered Medline: 19960829

A method for the identification of high-affinity ligands to SH2 domains by fluorescence-activated bead sorting (FABS) was established.

Recombinant SH2 domains, expressed as glutathione S-transferase (GST) fusion proteins, were incubated with a phosphotyrosine (Y\*)-containing peptide library. 6.4 x 10(5) individual peptides of nine amino acids in length (EPX6Y\*X19X7X19X7X6) were each displayed on beads. Phosphopeptide interaction of a given SH2 domain was monitored by binding of fluorescein isothiocyanate-labeled antibodies directed against GST. High-fluorescence beads were isolated by flow cytometric sorting. Subsequent pool sequencing of the selected beads revealed a distinct pattern of phosphotyrosine-containing motifs for each individual SH2 domain: the SH2 domain of the adapter protein Grb2 predominantly selected beads with the sequence Y\*ENDP, whereas the C-terminal SH2 domain of the tyrosine kinase Syk selected Y\*EELD, each motif representing the most frequently found residues C-terminal to the phosphotyrosine. For deconvolution studies, soluble phosphotyperides comprising variations of the Grb2 motifs were resynthesized and analyzed by surface plasmon resonance.

L5 ANSWER 10 OF 66 MEDLINE

ACCESSION NUMBER: 96234073 MEDLINE

DOCUMENT NUMBER: 96234073 PubMed ID: 8650202

Molecular cloning of the Golgi apparatus uridine diphosphate.N-acetylglucosamine transporter from Kluyveromyces lactis.

AUTHOR: Abeijon C; Robbins P W; Hirschberg C B

Department of Biochemistry and Molecular Biology, University of Massachusetts Medical Center, Worcester, 01655, USA.

CONTRACT NUMBER: GM 30365 (NIGMS)
GM 45188 (NIGMS)
GM 45188 (NIGMS)
FROCEEDINGS OF THE NATIONAL ACADEMY OF SCIENCES OF THE

SOURCE:

OF 45166 (HIGHS) PROCEEDINGS OF THE NATIONAL ACADEMY OF SCIENCES OF THE UNITED STATES OF AMERICA, (1996 Jun 11) 93 (12) 5963-8. Journal code: PV3; 7505876. ISSN: 0027-8424. United States

PUB. COUNTRY:

Journal; Article; (JOURNAL ARTICLE)

LANGUAGE:

English Priority Journals GENBANK-U48413 FILE SEGMENT:

OTHER SOURCE: ENTRY MONTH: 199607

ENTRY DATE:

Entered STN: 19960805 Last Updated on STN: 19960805

Entered Medline: 19960725

The mannan chains of Kluyveromyces lactis mannoproteins are similar to those of Saccharomyces cerevisiae except that they lack mannose phosphate and have terminal alpha1->2-linked N-acetylglucosamine. The biosynthesis of these chains probably occurs in the lumen of the Golgi apparatus, by analogy to S. cerevisiae. The sugar donors, GDP-mannose and UDP-GloNac, must first be transported from the cytosol, their site of synthesis, via specific Golgi membrane transporters into the lumen where they are substrates in the biosynthesis of these mannoproteins. A mutant of K. lactis, mnn2-2, that lacks terminal N-acetylglucosamine in its mannan chains in vivo, has recently been characterized and shown to have a specific defect in transport of UDP-GloNac into the lumen of Golgi vesicles in vitro. We have now cloned the gene encoding the K. lactis Golgi membrane UDP-GloNac transporter by complementation of the mnn2-2 mutantion. The mnn2-2 mutant was transformed with a genomic library from wild-type K. lactis in a pKD1-derived vector; transformants were isolated and phenotypic correction was monitored following cell surface labeling with fluorescein isothiocyanate conjugated to Griffonia simplicifolia II lectin, which binds terminal N-acetylglucosamine, and a fluorescent activated cell sorter. A 2.4-kb DNA fragment was found to restore the wild-type lectin binding phenotype. Upon loss of the The mannan chains of Kluyveromyces lactis mannoproteins are similar to to restore the wild-type lectin binding phenotype. Upon loss of the plasmid containing this fragment, reversion to the mutant phenotype occurred. The above fragment contained an open reading frame for a multitransmembrane spanning protein of 328 amino acids. The protein contains a leucine zipper motif and has high homology to predicted

proteins from S. cerevisiae and C. elegans. In an assay in vitro, Golgi vesicles isolated from the transformant had regained their ability to transport UDP-GlcNAc. Taken together, the above results strongly suggest that the cloned gene encodes the Golgi UDP-GlcNAc transporter of K.

=> FIL STNGUIDE COST IN U.S. DOLLARS SINCE FILE TOTAL ENTRY SESSION 33.87 FULL ESTIMATED COST 28.91

SINCE FILE DISCOUNT AMOUNTS (FOR QUALIFYING ACCOUNTS) TOTAL ENTRY SESSION CA SUBSCRIBER PRICE

FILE 'STNGUIDE' ENTERED AT 08:34:01 ON 03 MAY 2002 USE IS SUBJECT TO THE TERMS OF YOUR CUSTOMER AGREEMENT COPYRIGHT (C) 2002 AMERICAN CHEMICAL SOCIETY, JAPAN SCIENCE AND TECHNOLOGY CORPORATION, AND FACHINFORMATIONSZENTRUM KARLSRUHE

FILE CONTAINS CURRENT INFORMATION LAST RELOADED: Apr 26, 2002 (20020426/UP).

d 11- ibib abs

YOU HAVE REQUESTED DATA FROM FILE 'MEDLINE, AGRICOLA, CAPLUS, EMBASE' - CONTINUE? (Y) /N:v

YOU HAVE REQUESTED DATA FROM 56 ANSWERS - CONTINUE? Y/(N):y

ANSWER 11 OF 66 EMBASE COPYRIGHT 2002 ELSEVIER SCI. B.V. SSION NUMBER: 96343366 EMBASE

ACCESSION NUMBER: 1996343366

DOCUMENT NUMBER:

AUTHOR:

1996143366
Large DNA fragment sizing by flow
cytometry: Application to the characterization of
Pl artificial chromosome (PAC) clones.
Huang Z.; Petty J.T.; O'Quinn B.; Longmire J.L.; Brown
N.C.; Jett J.H.; Keller R.A.
Chemical Science Technology Division, Los Alamos National
Laboratory, Los Alamos, NM 87545, United States
Nucleic Acids Research, (1996) 24/21 (4202-4209).
ISSN: 0305-1048 CODEN: NARHAD CORPORATE SOURCE:

COUNTRY: DOCUMENT TYPE:

United Kingdom
Journal; Article
029 Clinical Biochemistry
English FILE SEGMENT:

SUMMARY LANGUAGE:

ULY Clinical Biochemistry
UNAGE: English
Aflow cytometry-based, ultrasensitive
fluorescence detection technique is used to size individual DNA
fragments up to 167 kb in length. Application of this technology to the
sizing of Pl artificial chromosomes (PACs) in both linear and supercoiled
forms is described. It is demonstrated that this method is well suited to
characterizing PAC/BAC clones and will be very useful for the analysis of
large insert libraries. Fluorescence bursts are recorded as
individual, dye stained DNA fragments pass through a low power, focused,
continuous laser beam. The magnitudes of the fluorescence bursts
are linearly proportional to the lengths of the DNA fragments. The
histograms of the burst sizes are generated in < 3 min with < 1 pg of DNA.
Results on linear fragments are consistent with those obtained by
pulsed-field gel electrophoresis. In comparison with pulsed-field gel
electrophoresis, sizing of large DNA fragments by this approach is more
accurate, much faster, requires much less DNA, and is independent of the

L5 ANSWER 12 OF 66 ACCESSION NUMBER:

DOCUMENT NUMBER:

TITLE:

97084577 MEDLINE DUPLICATE 5
97084577 MEDLINE
97084577 PubMed ID: 8930920
Bacterial genetics by flow cytometry:
rapid isolation of Salmonella typhimurium acid-inducible
promoters by differential fluorescence induction.
Valdivia R H; Falkow S
Department of Microbiology and Immunology, Stanford
University School of Medicine, California 94305, USA..
valdivia@cmgm.stanford.edu
AI 26195 (NIAID)
DK 38707 (NIDDK)
MOLECULAR MICROBIOLOGY, (1996 Oct) 22 (2) 367-78.
Journal code: MOM; 8712028. ISSN: 0950-382X.
EMGLAND: United Kingdom
Journal; Article; (JOURNAL ARTICLE)
English

AUTHOR:

CORPORATE SOURCE:

CONTRACT NUMBER:

SOURCE:

PUB. COUNTRY:

LANGUAGE:

FILE SEGMENT:

OTHER SOURCE:

Priority Journals
GENBANK-U62708; GENBANK-U62709; GENBANK-U62710;
GENBANK-U62711; GENBANK-U62712; GENBANK-U62713;

GENBANK-U62714

199702 Entered STN: 19970306 ENTRY MONTH:

Y DATE: Intered STN: 19970306

Last Updated on STN: 19980206

Entered Medline: 19970225

The ability of Salmonella typhimurium to survive and replicate within murine macrophages is dependent on a low phagosomal pR. This requirement for an acidic vacuole suggests that low pH is an important environmental stimulus for the transcription of genes necessary for intracellular survival. To study the behaviour of acid-inducible genes in response to the phagosomal environment, we have applied a novel enrichment strategy, termed differential fluorescence induction (DFI), to screen an S. typhimurium library for promoters that are upregulated at pH 4.5. DFI utilizes a fluorescence-enhanced green fluorescent protein (GFP) and a fluorescence-activated cell sorter (FACS) to perform genetic selection. In the presence of an inducing stimulus, such as low pH, a PACS is used to sort highly fluorescent bacterial clones bearing random promoters fused to the mutant GFP protein (GFPmut). This population is then amplified at neutral pH and the least fluorescent population is sorted. Sequential

ANSWER 13 OF 66 MEDLINE SION NUMBER: 97129113

ACCESSION NUMBER:

DOCUMENT NUMBER: TITLE:

sorts for fluorescent and non-fluorescent bacteria in the presence or absence of inducing conditions rapidly enriches for promoter fusions that are regulated by the inducing stimulus. We have identified eight acid-inducible promoters and quantified their expression in response to pH 4.5 and to the phagosome milieu. These acid-inducible promoters exhibited extensive homology to promoter regions of genes encoding for cell-surface-maintenance enzymes, stress proteins, and generalized efflux pumps. Only a subset of these promoters was induced in macrophages with kinetics and levels of expression that do not necessarily correlate with in vitro pH-shock induction. This suggests that while low pH is a relevant inducer of intracellular gene expression, additional stimuli in the macrophage can modulate the expression of acid-inducible genes. sorts for fluorescent and non-fluorescent bacteria in genes.

MEDLINE

```
97129113 PubMed ID: 8973653
Isolation and characterization of a 14.5-kDa
trichloroacetic-acid-soluble translational inhibitor
protein from human monocytes that is upregulated upon
                                                                                                 cellular differentiation
                                                                                               cellular differentiation.
Schmiedeknecht G; Kerkhoff C; Orso E; Stohr J; Aslanidis C;
Nagy G M; Knuechel R; Schmitz G
Institute of Clinical Chemistry and Laboratory Medicine,
University of Regensburg, Germany.
EUROPEAN JOURNAL OF BIOCHEMISTRY, (1996 Dec 1) 242 (2)
  AUTHOR:
  CORPORATE SOURCE:
  SOURCE:
                                                                                                Journal code: EMZ; 0107600. ISSN: 0014-2956. GERMANY: Germany, Federal Republic of Journal; Article; (JOURNAL ARTICLE)
  PUB. COUNTRY:
 LANGUAGE:
FILE SEGMENT:
                                                                                                 Priority Journals
OTHER SOURCE:
ENTRY MONTH:
                                                                                                 GENBANK-X95384
                                                                                                 199701
                                                                                                 Entered STN: 19970219
  ENTRY DATE:
                 Last Updated on STN: 19970219
Entered Medline: 19970128
A trichloroacetic-acid-soluble 14.5-kDa protein (p14.5) has been isolated from human mononuclear phagocytes (MMP) by a combination of trichloroacetic acid extraction, preparative electrophoresis and hydrophobic affinity chromatography; five tryptic peptides were subjected to protein sequencing. The full-length CDNA of the protein was cloned and sequenced from a lambda gtl1 human liver library. The CDNA showed a remarkable similarity to a rat protein preferentially expressed in hepatocytes and renal tubular epithelial cells. The encoded protein is 137 amino acids long and similar to members of a new hypothetical family of small proteins with presently unknown function, named YER057c/YJGF. Human recombinant p14.5 inhibits in vitro protein synthesis in a rabbit reticulocyte lysate system. Unlike other inhibitors of protein synthesis, p14.5 is not phosphorylated despite the presence of putative phosphorylation sites. The p14.5 mRNA is weakly expressed in freshly isolated monocytes but is significantly upregulated when these monocytes are subjected to differentiation. This is also reflected by a differentiation-dependent increase in the protein concentration as demonstrated by immunoblots from cytosolic fractions and fluorescence-activated flow cytometry of permeabilized cells. A differentiation-dependent mRNA and protein expression in a variety of liver and kidney tumor cells and a high expression in a variety of liver and kidney tumor cells and a high expression in a variety of liver and kidney tumor cells and only weak expression was found in other human tissues as evaluated by northern blot analysis. The preferential localization of the immunoreaction product seemed to be cytoplasmatic but, in less differentiated cells and only weak expression was found in other human tissues as evaluated by northern blot analysis. The preferential localization of the immunoreaction product seemed to be cytoplasmatic but, in less differentiated cells and only to the protein
                                                                                               Last Updated on STN: 19970219
Entered Medline: 19970128
                       ANSWER 14 OF 66
                                                                                                                      MEDLINE
  ACCESSION NUMBER:
                                                                                              96200387
96200387
                                                                                                                                                           MEDLINE
                                                                                                 96200387 MEDLINE
96200387 PubMed ID: 8927696
Cytogenetic damage and the radiation-induced G1-phase
 DOCUMENT NUMBER:
 TITLE:
                                                                                                 checkpoint.
                                                                                               Gupta N; Vij R; Haas-Kogan D A; Israel M A; Deen D F;
Morgan W F
AUTHOR .
                                                                                             Morgan W F
Brain Tumor Research Center of the Department of
Neurological Surgery, University of California San
Francisco 94143, USA.
CA13525 (NCI)
RADIATION RESEARCH, (1996 Mar) 145 (3) 289-98.
Journal code: QMP; 0401245. ISSN: 0033-7587.
United States
CORPORATE SOURCE:
CONTRACT NUMBER:
 SOURCE:
PUB. COUNTRY:
                                                                                                Journal; Article; (JOURNAL ARTICLE)
LANGUAGE .
                                                                                                English
FILE SEGMENT:
ENTRY MONTH:
                                                                                               Priority Journals
                    Y MONTH: 199611
Y DATE: Entered STN: 19961219
Last Updated on STN: 19961219
Entered Medline: 19961114
It is proposed that genomic integrity is preserved after DNA damage in a variety of ways. X irradiation induces a p53-dependent G1-phase cell cycle checkpoint which putatively allows time for repair of DNA damage. The p53 protein is also involved in the initiation of apoptosis after radiation-induced DNA damage, presumably leading to the elimination of lethally damaged cells from the irradiated population. To test the hypothesis that repair occurs in the additional time provided by the activation of the G1-phase checkpoint, we investigated whether the presence of a G1-phase arrest modified the frequency and type of chromosomal rearrangements at the first mitosis after irradiation.
                                                                                               199611
```

Isogenic cell lines derived from the same human glioma cell line, but differing in p53 status, were used. Purified G1-phase cells, isolated by centrifugal elutriation and X-irradiated, were studied. The wild-type p53 cell line demonstrated a dose-dependent arrest during G1 phase, as determined by flow cytometry. These cells remained in G1-phase as long as 48 h after irradiation. Cells expressing a dominant-negative p53 mutation accumulated to a much lesser extent in G1 phase after irradiation. Cells lacking the G1-phase checkpoint showed increased survival at all radiation doses. There were no significant differences in the type or frequency of total chromosomal aberrations in mitotic cells from either cell line after 1,2,4 or 6 Gy X rays, as measured by conventional cytogenetic analysis. There was an increase, however, in the number of reciprocal translocations in mitotic cells with mutant p53 (lacking a G1-phase checkpoint), as measured by fluorescence in situ hybridization with a chromosome 4-specific DNA library, but only after 6 Gy. The results suggest that the presence of a well-defined p53-dependent G1-phase arrest does not reduce chromosomal aberrations caused by low doses of ionizing radiation markedly, but may reduce the overall degree of survival by triggering other G1-phase events. Isogenic cell lines derived from the same human glioma cell line, but

ANSWER 15 OF 66 MEDLINE

ACCESSION NUMBER: DOCUMENT NUMBER:

97130619 MEDLINE 97130619 PubMed ID: 8976382

TITLE:

Orline Pubmed ID: 8976382
Characterization of a flow-sorted human chromosome 10
cosmid library by FISH.
Ma N S; Zheng C; Benchekroun Y; Deaven L L; Longmire J L;
Moir D T; Mao J
Department of Human and Molecular Genetics, Genome AUTHOR:

CORPORATE SOURCE:

Therapeutics Corporation, Waltham, USA. R01-HG00122 (NHGRI) R44-HG00491 (NHGRI)

CONTRACT NUMBER:

CYTOGENETICS AND CELL GENETICS, (1996) 74 (4) 266-71.
Journal code: DXK; 0367735. ISSN: 0301-0171.
Switzerland SOURCE:

PUB. COUNTRY:

Journal; Article; (JOURNAL ARTICLE)

LANGUAGE: English FILE SEGMENT:

Priority Journals GDB-G00631948 OTHER SOURCE: ENTRY MONTH: 199701 Entered STN: 19970128 ENTRY DATE:

Last Updated on STN: 19980206 Entered Medline: 19970115

Entered Medline: 19970115
Fluoresence in situ hybridization (FISH) was used to localize cosmids to regions of human chromosome 10. A total of 301 cosmids were selected randomly from a flow-sorted human chromosome 10 cosmid library constructed from human x hamster cell line 762-8A and arrayed in microtiter storage dishes. Over 70% (211/301) of the cosmids mapped to unique regions of chromosome 10. About 7% (22/301) produced multiple hybridization signals indicative of chimeric clones or sequences repeated at low copy number. Three cosmids (3/301, or 1%) hybridized to the centromeric regions of chromosome 10 and one or more other human chromosomes. About 19% (59/301) consisted mostly or entirely of hamster DNA inserts, and about 2% (6/301) appeared to be nonrecombinants.

ANSWER 16 OF 66 MEDLINE

ACCESSION NUMBER: 1998040649 MEDLINE DOCUMENT NUMBER:

1998040649 MEDLINE
98040649 PubMed ID: 9373318
Separation of E. coli expressing functional cell-wall bound antibody fragments by FACS.
Fuchs P; Weichel W; Dubel S; Breitling F; Little M
Recombinant Antibody Research Group, Deutsches
Krebsforschungszentrum, Heidelberg, Germany.
IMMUNOTECHNOLOGY, (1996 Jun) 2 (2) 97-102.
Journal code: CRO; 9511979. ISSN: 1380-2933. AUTHOR: CORPORATE SOURCE: SOURCE:

PUB. COUNTRY:

Journal; Article; (JOURNAL ARTICLE)

English Priority Journals FILE SEGMENT:

LANGUAGE:

ENTRY MONTH: 199712 ENTRY DATE:

ANSWER 17 OF 66 CAPLUS COPYRIGHT 2002 ACS ACCESSION NUMBER:

TITLE:

PLUS COPYRIGHT 2002 ACS 1996:217232 CAPLUS Flow cytometric screening of antibody fragment contact residue libraries displayed at the surface of escherichia coli: A model system for the development of bacterial surface display for antibody selection.

```
08/876,276 Search Strategy/Results
                                                                                                              Daugherty, Patrick; Chen, Gang; Iverson, Brent;
Georgiou, George
Departments Chemical Engineering, University Texas,
Austin, TX, 78712, USA
Book of Abstracts, 211th ACS National Meeting, New
Orleans, LA, March 24-28 (1996), BIOT-076. American
Chemical Society: Washington, D. C.
   AUTHOR (S)
   CORPORATE SOURCE:
   SOURCE:
                                                                                                                CODEN: 62PIAJ
                   CODEN: 62PIAJ

MENT TYPE: Conference; Meeting Abstract

SUAGE: English

The design of antibodies and peptides for particular applications has been made possible in recent years through the advent of various

1thrary generation and screening technologies. In particular, the display of libraries of antibody fragments on the external surface of filamentous phage as fusions to major coat proteins has yielded antibody fragments and peptides for numerous applications. However, the screening of antibody libraries by phage display has proven to be technol. challenging and suffers from certain limitations. Recently, we have shown that single-chain antibodies (serV) may be displayed at the surface of Escherichia coli, and subsequently that clones as rare as one in 105 may be selected from cells displaying non-antigen binding proteins. In order to demonstrate the potential of bacterial surface display in conjunction with fluorescence activated cell sorting (PACS), for high-affinity antibody fragment selection, we have constructed small antibody libraries displayed at the surface of Escherichia coli, and screened them for high-affinity antigen binding. Several plasmid libraries encoding surface-displayed single-chain antibody fragments have been generated by PCR randomization of key antigen-contact residues in the digoxin scFv, as suggested by the known crystal structure. The plasmid libraries were transformed into Escherichia coli and screened for high affinity antigen binding by incubation of library aliquots with
                                                                                                               Conference; Meeting Abstract
   DOCUMENT TYPE:
   LANGUAGE:
                     libraries were transformed into Escherichia coli and screened for high affinity antigen binding by incubation of library aliquots with a fluorescent antigen conjugate followed by FACS. Amplification of the sorted populations yielded populations enriched for high affinity antigen binding, as detd. by flow cytometry and DNA sequence anal. of individual clones. Several important parameters for bacterial surface display, library construction, and FACS screening have been analyzed and will be presented.
L5 ANSWER 18 OF 66 MEDLINE
ACCESSION NUMBER: 97150954
DOCUMENT NUMBER: 97150954 Pu
                                                                                                                                               MEDLINE
                                                                                        97150954 MEDLINE
97150954 PubMed ID: 8995490
Development of DNA libraries specific for Chinese hamster
chromosomes 3, 4, 9, 10, X, and Y by DOP-PCR.
Xiao Y; Slijepcevic P; Arkesteijn G; Darroudi F; Natarajan
DOCUMENT NUMBER:
AUTHOR:
 CORPORATE SOURCE:
                                                                                         MGC-Department of Radiation Genetics and Chemical
                                                                                        Mutagenesis, Leiden University, The Netherlands.
CYTOGENETICS AND CELL GENETICS, (1996) 75 (1) 57-62.
Journal code: DXK; 0367735. ISSN: 0301-0171.
 SOURCE:
                                                                                         Switzerland
PUB. COUNTRY:
                                                                                          Journal; Article; (JOURNAL ARTICLE)
LANGUAGE:
                                                                                        English
FILE SEGMENT:
ENTRY MONTH:
                                                                                        Priority Journals
199701
                                                                                        Entered STN: 19970219
Last Updated on STN: 19970219
ENTRY DATE:
                    Last Updated on STN: 19970219
Entered Medline: 19970130
In a previous study of ours, DNA libraries specific for Chinese hamster chromosomes 1, 2, 3, 5, 6, 7, and 8 were constructed by single laser flow sorting and the linker adapter PCR method. Since a single laser flow sorting of chromosomes is based on chromosome size, it is difficult to separate chromosomes of equal size, such as chromosome 3 and the X chromosome. This problem can be circumvented by bivariate flow sorting. In the present study, we have used bivariate flow sorting of Chinese hamster chromosomes and the degenerate oligonucleotide-primed polymerase chain reaction (DOP-PCR) for generating DNA libraries specifically for chromosomes 3, 4, 9, 10, X, and Y. Pluorascence in situ hybridization (PISH) analyses indicate that the probe libraries are highly specific and can be applied to various cytogenetic studies.
                      specific and can be applied to various cytogenetic studies.
                    ANSWER 19 OF 66
                                                                                                             MEDLINE
                                                                                      96305137
96305137
                                                                                                                                    MEDLINE
PubMed ID: 8707053
 ACCESSION NUMBER.
 DOCUMENT NUMBER:
                                                                                      PAGS-optimized mutants of the green fluorescent protein (GFP).

Cormack B P; Valdivia R H; Falkow S
Department of Microbiology and Immunology, Stanford University School of Medicine, CA 94305-5402, USA.
AUTHOR:
CORPORATE SOURCE:
                                                                                      COTMACK@Cmgm.stanford.edu
AI 36396 (NIAID)
GENE, (1996) 173 (1 Spec No) 33-8.
Journal code: FOP; 7706761. ISSN: 0378-1119.
 CONTRACT NUMBER:
SOURCE:
PUB. COUNTRY:
                                                                                        Netherlands
                                                                                        Journal; Article; (JOURNAL ARTICLE)
                                                                                        English
                                                                                        Priority Journals
GENBANK-U73901; GENBANK-U89685; GENBANK-U89686
FILE SEGMENT:
OTHER SOURCE:
ENTRY MONTH:
                   Y MONTH: 199609
Y MONTH: 199609
Y MONTH: 199609
Y DATE: Entered STN: 19960919
Last Updated on STN: 19980206
Entered Medline: 19960911
We have constructed a library in Escherichia coli of mutant gfp
genes (encoding green fluorescent protein, GFP) expressed from a
tightly regulated inducible promoter. We introduced random amino acid (aa)
substitutions in the twenty aa flanking the chromophore Ser-Tyr-Gly
sequence at aa 65-67. We then used fluorescence-activated cell
sorting (FACS) to select variants of GFP that fluoresce between 20-and
35-fold more intensely than wild type (wt), when excited at 488 mm.
Sequence analysis reveals three classes of aa substitutions in GFP. All
three classes of mutant proteins have highly shifted excitation maxima. In
addition, when produced in E. coli, the folding of the mutant proteins is
more efficient than folding of wt GFP. These two properties contribute to
a greatly increased (100-fold) fluorescence intensity, making
the mutants useful for a number of applications.
                                                                                        199609
ENTRY DATE:
```

L5 ANSWER 20 OF 66 ACCESSION NUMBER: MEDLINE

97381297 MEDLINE 97381297 PubMed ID: 9238629 DOCUMENT NUMBER:

AUTHOR

CORPORATE SOURCE:

CONTRACT NUMBER:

97381297 PubMed ID. 9238629
The use of a combinatorial library method to isolate human tumor cell adhesion peptides.
Pennington M E; Lam K S; Cress A E
Department of Radiation Oncology, University of Arizona, Tucson 85824, USA.
CA 17094 (NCI)
CA 23047 (NCI)
CA 56666 (NCI)
MOLECULAR DIVERSITY, (1996 Oct) 2 (1-2) 19-28.
Journal code: CWL; 9516534. ISSN: 1381-1991.
Netherlands SOURCE:

Netherlands PUB. COUNTRY: Journal; Article; (JOURNAL ARTICLE)

English LANGUAGE:

FILE SEGMENT: Priority Journals ENTRY MONTH: ENTRY DATE: 199708 Entered STN: 19970902

Type Table : Entered STN: 19970902

Last Updated on STN: 19970902

Entered Meddine: 19970821

Tumor cell progression is dependent in part on the successful adhesive interactions of the cells with the extracellular matrix. In this study, a new approach is described to isolate linear peptide ligand candidates involved in cellular adhesion. A synthetic combinatorial peptide library based on the 'one-bead-one-peptide' concept was incubated with live human prostate cancer cells for 90 min at 37 degrees C. The peptide bead coated with a monolayer of cells was then isolated for microsequencing. The DU145 (DU-H) cells were chosen since-they have been previously characterized as containing elevated levels of a laminin receptor for cell adhesion, the alpha 6 beta 1 integrin on the cell surface. The use of a function-blocking antibody (GoH3) allows for the detection of peptides which are alpha 6-specific ligand candidates. From two different libraries (linear 9-mer and 11-mer) of a total of 1,500,000 beads, 68 peptide beads containing attached cells were isolated. These positive beads were then retested to determine the ability of the GoH3 antibody to block binding of the cells to the peptide beads. The alpha 6 beads, 68 peptide beads containing attached cells were isolated. These positive beads were then retested to determine the ability of the GoH3 antibody to block binding of the cells to the peptide beads. The alpha 6 integrin candidate peptide beads (five in total) were recovered and two of the beads were microsequenced. These two peptides, RU-1 (LNIVS-VNGRHX) and RX-1 (DNRIRLOAKXX), resemble the previously reported active peptide sequences (GD-2 and AG-73) from native laminin. The RU-1, RX-1 and AG-73 peptides were tested for their ability to support cell attachment and to bind the cell surface of DU-H prostate carcinoma cells in suspension using fluorescence-activated cell-sorting (FACS) analysis. Both RU-1 and AG-73 peptides supported cellular attachment within 1 h. In contrast, after 1 h, EHS laminin supported both cellular attachment and spreading. The RX-1 peptide exhibited only weak binding to the DU-H prostate carcinoma cells. FACS analysis indicated that AG-73 peptide attached to tumor cell surfaces over a range of concentrations, whereas the RU-1 peptide showed a homogeneous concentration required for attachment. The described strategy for screening a random peptide library offers three advantages: (1) ligands for conformationally sensitive receptors of adhesion can be isolated using live cells; (ii) specific binding can be selected for using function-blocking antibodies; and (iii) peptides supporting adhesion independent of spreading properties can be distinguished. In principle, specific adhesive peptides without prior knowledge of the sequence could be isolated for any epithelial cell surface receptor for which a function-blocking reagent is available.

ANSWER 21 OF 66 MEDLINE

ACCESSION NUMBER: DOCUMENT NUMBER: 95399346 95399346 MEDLINE PubMed ID: 7669720

95399346 PubMed ID: 7669720
Positive selection of growth-inhibitory genes.
Maines J Z; Sunnarborg A; Rogers L M; Mandavilli A;
Spielmann R; Boyd F T
Department of Laboratory Medicine and Pathology, University
of Minnesota, Minneapolis 55455, USA.
CA-56967 (NCI)
CELL GROWTH AND DIFFERENTIATION, (1995 Jun) 6 (6) 665-71.
Journal code: AYH; 9100024. ISSN: 1044-9523.
United States
Journal: Article: (JOURNAL ARTICLE) TITLE: AUTHOR:

CORPORATE SOURCE:

CONTRACT NUMBER:

SOURCE:

PUB. COUNTRY:

Journal; Article; (JOURNAL ARTICLE)

LANCHAGE .

English Priority Journals FILE SEGMENT: ENTRY MONTH: 199510 Entered STN: 19951026

NONTH: 199510

YOUNTE: Entered STN: 19951026

Entered Medline: 19951019

We have isolated a limited set of cDNAs that limit cell proliferation using a unique assay based on the dilution of a lipophilic fluorescent dye as transfected cells divide. The identification of growth-inhibitory factors has been limited by the lack of a strong assay for growth inhibitors. A growth-inhibited cell does not grow and so is at a selective disadvantage in vitro when compared with any growing cell. Several assays have been used to screen for growth-inhibitory genes; however, these approaches are either very difficult to implement, leaky, or not comprehensive. We have developed an assay that selects for cDNAs capable of inhibiting proliferation in which cells are nonspecifically labeled with a lipophilic fluorescent dye, PKH-2, and subsequently transfected with a cDNA library made from growth-inhibited cells. With each cell division, the amount of dye per cell is reduced by one-half. Over time, growth-inhibited cells will retain more dye per cell relative to actively growing cells. The population is then analyzed by fluorescence-activated cell sorting, and the brightest cells in the population are isolated. This assay has allowed us to select pools of cDNAs enriched for growth-inhibitory activities and may provide a general method for identifying growth-inhibitory genes active in varying biological contexts. We report here the successful application of the dye retention assay to the selection of cDNAs that inhibit epithelial cell proliferation. cell proliferation.

```
08/876,276 Search Strategy/Results
                                                                                                                                                   PubMed ID: 7731047
                                                                                                  95248551
  DOCUMENT NUMBER:
                                                                                                95248551 PubMed ID: 7731047
Selection and application of human single chain Fv antibody fragments from a semi-synthetic phage antibody display library with designed CDR3 regions.
de Kruif J; Boel E; Logtenberg T
Department of Immunology, University of Utrecht, The Netherlands.
JOURNAL OF MOLECULAR BIOLOGY, (1995 Apr 21) 248 (1) 97-105.
JOURNAL Oct BOELOGIAR BIOLOGY, COMPANDED COMPAND
  AUTHOR:
 CORPORATE SOURCE:
  SOURCE:
  PUB. COUNTRY:
                                                                                                  Journal; Article; (JOURNAL ARTICLE)
                                                                                                     English
         ANGUAGE:
                                                                                                  Priority Journals
GENBANK-X83616; GENBANK-X83712; GENBANK-X83713;
GENBANK-X83714
  FILE SEGMENT:
  OTHER SOURCE
 ENTRY MONTH:
ENTRY DATE:
                                                                                                   199505
                     Y MONTH: 199505
Y DATE: Entered STN: 19950608
Last Updated on STN: 19960129
Entered Medline: 19950526
We have constructed a large (3.6 x 10(8) clones) phage display
library of human single chain Fv (scFv) antibody fragments by
combining 49 germline VH genes with synthetic heavy chain CDR3 (HCDR3)
regions and seven light chains. The HCDR3 regions varied in length between
6 and 15 residues and were designed to contain fully randomized stretches
of amino acid residues flanked by regions of limited residue variability
that were composed of amino acid residues that frequently occur in natural
antibodies. We reasoned that this approach would increase the frequency of
functional molecules in our library and, in addition, permit us
to efficiently utilize available cloning space. By direct selection on
solid phase-bound antigens we obtained phage antibodies with binding
activities to 13 different antigens, including Von Willebrand factor, the
DNA-binding HMG box of transcription factor TCP-1 and the tumor antigen
EGP-2. In addition, we applied a competitive selection procedure to target
phage antibodies to the desired portion of a recombinant fusion protein
and to select phage antibodies capable of discriminating between the two
highly homologous homeobox proteins PBXIa and PBX2. The functional
capacity of monoclonal phage antibodies was assessed in
                                                                                                   Entered STN: 19950608
                        and the state of monoclonal phage antibodies was assessed in immuno-histochemical staining of tissue specimens. Western blotting assays and immunofluorescent analysis of cells by flow cytometry. The results demonstrate that this large human phage antibody library contains a broad assortment of binding
                         specificities that can be applied in a variety of biochemical assays.
                       ANSWER 23 OF 66
                                                                                                                         MEDLINE
 ACCESSION NUMBER:
DOCUMENT NUMBER:
                                                                                                96159517 MEDLINE
96159517 PubMed ID: 8574897
                                                                                                  Characterization of a human chromosome 22 enriched bacterial artificial chromosome sublibrary.
 TITLE:
                                                                                                Journal code: CEK; 9509403.
AUTHOR:
CORPORATE SOURCE:
 SOURCE:
                                                                                                  Netherlands
  PUB. COUNTRY:
                                                                                                  Journal; Article; (JOURNAL ARTICLE)
LANGUAGE:
FILE SEGMENT:
ENTRY MONTH:
ENTRY DATE:
                                                                                                   English
                                                                                                  Priority Journals
                                                                                                 199603
Entered STN: 19960321
                       Last Updated on STN: 19960321
Entered Medline: 19960312
Selection of chromosomal sublibraries from total human genomic libraries
                     Selection of chromosomal sublibraries from total human genomic libraries is critical for chromosome-based physical mapping approaches. We have previously reported a method of screening total human genomic library using flow sorted chromosomal DNA as a hybridization probe and selection of a human chromosome 22-enriched sublibrary from a total human bacterial artificial chromosome (BAC) library (Nucleic Acids Res 1995; 23: 1838-39). We describe here further details of the method of construction as well as characterization of the chromosome 22-enriched sublibrary thus constructed. Nearly 40% of the BAC clones that have been mapped by fluorescence in situ hybridization (FISH) analysis were localized to chromosome 22. By screening the sublibrary using chromosome 22-specific hybridization probes, we estimated that the sublibrary represents at least 2.5 x coverage of chromosome 22. This is in good agreement with the results from FISH mapping experiments. FISH map data also indicate that chromosome 22-specific BACs in the sublibrary represent all the subregions of chromosome 22.
                         represent all the subregions of chromosome 22.
                       ANSWER 24 OF 66 AGRICOLA
ACCESSION NUMBER:
DOCUMENT NUMBER:
                                                                                                                           95:52968 AGRICOLA
CAT10701060
                                                                                                                         CAT10701060
Immunocytochemical methods and protocols.
Javois, Lorette C.
DNAL (QH506.M45 no.34)
94-22475
c1994 xii, 435 p.: ill.; 24 cm
Publisher: Totowa, N.J.: Humana Press, c1994.
Series: Methods in molecular biology (Clifton, N.J.); 34
TITLE:
 AUTHOR (S)
 AVAILABILITY:
LC CONTROL NO.:
SOURCE:
NOTE:
```

34. ISBN: 089603285X (acid-free paper).
Includes bibliographical references and index.
Antibody preparation -- Overview of antibody use in immunocytochemistry -- Purification of antibodies using ammonium sulfate fractionation or gel filtration -- Purification of antibodies using ion-exchange chromatography -- Purification of antibodies using affinity chromatography -- Purification of antibodies using protein A-sepharose and FPLC -- Conjugation of fluorochromes to antibodies -- Biotinylation of antibodies -- Tissue preparation for light microscopic analyses -- Overview of cell fixation and permeabilization.
Preparation of frozen sections for analysis --

```
Processing of cell-touch preparations -- Processing of cytological specimens -- Processing of tissue-culture cells -- Processing of tissue specimens -- Light microscopic detection systems -- Overview of fluorophores -- Direct immunofluorescent labeling of cells -- Fluorescence labeling of surface antigens of attached or suspended tissue-culture cells -- Fluorescence labeling of intracellular antigens of attached or suspended tissue-culture
                                                                                        antigens of attached or suspended tissue-culture
                                                                                     cells.

Fluorescent labeling of surface or
intracellular antigens in whole-amounts -- Overview of
antigen detection through enzymatic activity --
Peroxidase-antiperoxidase (PAP) method --
Avidin-biotin complex (ABC) method -- Labeled avidin
binding (LAB) method -- Avidin-biotin labeling of
cellular antigens in cryostat-sectioned tissue --
ELISA on attachment-dependent or suspension grown
cells -- Use of immunogold with silver enhancement --
Fluorescence-activated cell sorter (FACS)
analyses -- Overview of flow
cytometry and fluorescent probes for
                                                                                        cytometry and fluorescent probes for
                                                                                     cytometry.
Tissue disaggregation -- Indirect immunofluorescent
labeling of viable cells -- Indirect immunofluorescent
labeling of fixed cells -- Fluorescent
labeling of DNA -- Deparaffinization and processing of
pathologic material -- Assay for phagocytosis -- Assay
for filamentous actin -- Assay for chemoattractant
binding -- Assay for oxidative metabolism -- Colloidal
gold detection systems for electron microscopic
analyses -- Fixation and embedding -- Preparation of
colloidal gold -- Conjugation of colloidal gold to
                                                                                        cytometry.
                                                                                        colloidal gold -- Conjugation of colloidal gold to
                                                                                        Colloidal gold/streptavidin methods -- Pre-embedding
                                                                                        labeling methods -- Postembedding labeling methods
Photography -- Overview of fluorescence
                                                                                     Photography -- Overview of fluorescence photomicrography -- Special applications -- Overview of fluorescence analysis with the confocal microscope -- Overview of laser microbeam applications as related to antibody targeting -- Nucleic acid immunocytochemistry: in situ hybridization to human chromosomes of an alkaline phosphatase-labeled centromeric probe -- Fluorescence in situ
                                                                                       hybridization using whole chromosome library
                                                                                       Overview of automated immunostainers -- Overview of immunocytochemical approaches to the differential diagnosis of tumors: diagnostic applications in the
                                                                                       clinical laboratory.
New Jersey; United States
PUB. COUNTRY:
DOCUMENT TYPE:
                                                                                       Bibliography; (MONOGRAPH)
U.S. Imprints not USDA, Experiment or Extension
English
FILE SEGMENT:
LANGUAGE:
                ANSWER 25 OF 66
                                                                                       MEDLINE
 ACCESSION NUMBER:
                                                                     94253194
                                                                                                                 MEDLINE
                                                                     94253194 MEDLINE
94253194 PubMed ID: 8195250
Expression cloning of a GM3-specific alpha-2,8-
sialyltransferase (GD3 synthase).
Sasaki K; Kurata K; Kojima N; Kurosawa N; Ohta S; Hanai N;
DOCUMENT NUMBER:
TITLE:
AUTHOR:
                                                                      Tsuji S; Nishi T
                                                                      Tokyo Research Laboratories, Kyowa Hakko Kogyo, Co., Ltd.,
CORPORATE SOURCE:
                                                                      Japan.

JOURNAL OF BIOLOGICAL CHEMISTRY, (1994 Jun 3) 269 (22)
SOURCE:
                                                                      15950-6.
                                                                      Journal code: HIV; 2985121R. ISSN: 0021-9258. United States
PUB. COUNTRY:
                                                                      Journal; Article; (JOURNAL ARTICLE)
                                                                      English
                                                                      Priority Journals
GENBANK-X77922
 FILE SEGMENT:
OTHER SOURCE:
ENTRY MONTH:
                                                                      199406
                                                                     Entered STN: 19940707
Last Updated on STN: 19980206
                Entered Medline: 19940630
A cDNA encoding a GM3-specific alpha-2,8-sialyltransferase (GD3 synthase) was obtained from an expression cDNA library of human melanoma cell line WM266-4 by enrichment of Namalwa KJM-1 cells highly expressing GD3 using an anti-GD3 antibody and a fluorescence-activated cell
               GD3 using an anti-GD3 antibody and a fluorescence-activated cell sorter. Selection of B-cell line Namalwa cells expressing transfected CDNAs in the presence of anti-GD3 monoclonal antibody KM641 gave a cDNA (pAMO-GD3) encoding a protein with a type II transmembrane topology as found for mammalian glycosyltransferases. The following evidence confirms that the cDNA encodes an alpha-2,8-sialyltransferase, which specifically converts GM3 to GD3. (i) Transfection of pAMO-GD3 into Namalwa KJM-1 cells leads to the appearance of GD3 and a GD3 synthase activity. (ii) Northern blot analysis revealed a correlation between the expression of this gene and GD3 in several cell lines. (iii) The putative COOH-terminal active domain of this cloned enzyme fused with protein A has been purified with IgG-Sepharose beads and has been shown to possess GD3-synthesizing activity, excluding the possibility that the cloned cDNA encodes a transacting factor inducing a GD3 synthase. The deduced primary sequence also contains the "sialyl motif" conserved among all the sialyltransferases cloned to date. The polymerase chain reaction analysis reveals that this gene is located on chromosome 12.
                ANSWER 26 OF 66
                                                                                      MEDLINE
                                                                                                                                                                                                   DUPLICATE 8
ACCESSION NUMBER:
DOCUMENT NUMBER:
                                                                    94286537
94286537
                                                                                                                  MEDLINE
                                                                                                        PubMed ID: 8016078
                                                                    Pure chromosome-specific PCR libraries from single sorted chromosomes.

VanDevanter D R; Choongkittaworn N M; Dyer K A; Aten J; Otto P; Behler C; Bryant E M; Rabinovitch P S
TITLE:
AUTHOR:
```

Tumor Institute, Swedish Medical Center, Seattle, WA 98104. . CORPORATE SOURCE: CONTRACT NUMBER:

AG 01751 (NIA) AG 10917 (NIA) DK 32971 (NIDDK)

PROCEEDINGS OF THE NATIONAL ACADEMY OF SCIENCES OF THE SOURCE:

UNITED STATES OF AMERICA, (1994 Jun 21) 91 (13) 5858-62. Journal code: PV3; 7505876. ISSN: 0027-8424.

PUB. COUNTRY: United States

Journal; Article; (JOURNAL ARTICLE)

LANGUAGE:

English Priority Journals 199407 FILE SEGMENT: ENTRY MONTH:

ENTRY DATE:

L5 ANSWER 27 OF 66 ACCESSION NUMBER: MEDLINE

94362249 MEDLINE

DOCUMENT NUMBER:

PubMed ID: 8080999
Rh(D) antigen expression and isolation of a new Rh(D) cDNA

Suyama K; Lunn R; Haller S; Goldstein J Lindsley P. Kimball Research Institute, New York Blood Center, New York 10021. BLOOD, (1994 Sep 15) 84 (6) 1975-81. Journal code: ABG; 7603509. ISSN: 0006-4971. AUTHOR: CORPORATE SOURCE:

SOURCE:

United States PUB. COUNTRY: Journal; Article; (JOURNAL ARTICLE)

LANGUAGE

FILE SEGMENT: Abridged Index Medicus Journals; Priority Journals

OTHER SOURCE: ENTRY MONTH: GENBANK-S73913 199410

ENTRY DATE:

Y MONTH: 199410
Y DATE: Entered STN: 19941021
Last Updated on STN: 19970203
Entered Medline: 19941012
Human erythroleukemic K562 cells are known to have several erythroid properties. K562 cells possess Rh mRNAs, but expression of Rh proteins has not previously been reported. We immunoprecipitated Rh protein from K562 cell lysate using rabbit anti-Rh and detected Rh(D) antigens on K562 cells using fluorescence-activated cell sorting (FACS). These results suggest that K562 cells will be useful as an expression model for most Rh antigens. We also cloned a new Rh(D) cDNA isoform (RhK562-II), from a K562 cDNA library using polymerase chain reaction (PCR) with 5' and 3' end oligonucleotides of the published Rh(e/E) antigen encoding cDNA sequence as primers. Sequence analysis showed that RhK562-II is composed 3' end oligonucleotides of the published Rh(e/E) antigen encoding cDNA sequence as primers. Sequence analysis showed that RhK562-II is composed of 951 nucleotides (316 amino acids), identical to the first 939 nucleotides (exons 1 to 6) of one of the Rh(D) cDNAs (RhXIII), except for nucleotide 654 (C-->G exchange). However, this exchange is the same as that of another published Rh(D) cDNA (RhPII cDNA). RhK562-II is deprived of exons 7 and 8 (nucleotides 940 to 1,153), followed by an identical sequence up to the 3' end of the open-reading frame of the RhXIII cDNA, which causes a frame-shift mutation and produces a premature stop codon. In vitro expression of RhK562-II using the transcription and translation rabbit reticulocyte lysate system produced two major Rh-related proteins (30 kD and 25 kD), which were immunoprecipitated by rabbit polyclonal anti-Rh and separated on sodium dodecyl sulfate-polyacrylamide gel electrophoresis (SDS-PAGE). electrophoresis (SDS-PAGE).

ANSWER 28 OF 66 MEDLINE

ACCESSION NUMBER: DOCUMENT NUMBER:

TITLE:

AUTHOR .

6 MEDLINE
95048345 MEDLINE
95048345 PubMed ID: 7959744
Generation of five high-complexity painting probe libraries from flow-sorted mouse chromosomes.
Weier H U; Polikoff D; Fawcett J J; Greulich K M; Lee K H; Cram S; Chapman V M, Gray J W
Division of Molecular Cytometry, School of Medicine,
Library of California San Prancisco 94143-0808

CORPORATE SOURCE:

University of California, San Francisco 94143-0808. RR01315 (NCRR) GENOMICS, (1994 Jun) 21 (3) 641-4. Journal code: GEN; 8800135. ISSN: 0888-7543. CONTRACT NUMBER: SOURCE:

PUB. COUNTRY: United States

Journal; Article; (JOURNAL ARTICLE) English

ANGUAGE FILE SEGMENT:

Priority Journals

ENTRY MONTH: ENTRY DATE:

Entered STN: 19950110
Last Updated on STN: 19950110
Entered Medline: 19941130

Mouse metaphase chromosomes were purified by flow sorting from the murine fibroblast cell line Mus spretus clone 5A. We sorted chromosomes that fell into five individual peaks based on the Hoechst 33258/chromomycin A3 DNA into five individual peaks based on the Hoechst 33258/chromomycin A3 DNA histogram: three peaks corresponding to the least amount of DNA and two peaks representing chromosomes with the most DNA content. This is the first example of the successful application of bivariate flow karyotyping to murine chromosome sorting. We then applied primer-directed in vitro DNA amplification using the polymerase chain reaction (PCR) to generate and label larger amounts of chromosome-specific DNA. In situ hybridization showed specific binding of the PCR products to mouse chromosomes Y, 19, 18, 3, and X as well as chromosomes 1 and 2. The combination of chromosome sorting from the M. spretus cell line and PCR proved to be highly valuable for generation of pools of DNA fragments that exhibit specific binding to mouse chromosomes and can be used to identify and delineate mouse metaphase chromosomes. metaphase chromosomes.

L5 ANSWER 29 OF 66 ACCESSION NUMBER: MEDLINE

DOCUMENT NUMBER:

94170829 MEDLINE 94170829 PubMed ID: 7510236 MRC OX19 recognizes the rat CD5 surface glycoprotein, but TITLE: does not provide evidence for a population of CD5bright B

AUTHOR: Vermeer L A; de Boer N K; Bucci C; Bos N A; Kroese F G;

CORPORATE SOURCE:

Department of Histology and Cell Biology, University of Groningen, The Netherlands. EUROPEAN JOURNAL OF IMMUNOLOGY, (1994 Mar) 24 (3) 585-92. SOURCE:

Journal code: EN5; 1273201. ISSN: 0014-2980. GERMANY: Germany, Federal Republic of Journal; Article; (JOURNAL ARTICLE)

PUB. COUNTRY:

LANGUAGE:

FILE SEGMENT: Priority Journals

OTHER SOURCE: ENTRY MONTH: GENBANK-X78985

199404

Entered STN: 19940420 ENTRY DATE:

Last Updated on STN: 19960129 Entered Medline: 19940412

Last Updated on STN: 19960129
Entered Medline: 19940412

To clone the rat CDS gene we first produced two rat CD5 probes. The probes were obtained by polymerase chain reaction (PCR) on rat genomic DNA using primers designed on conserved regions between mouse and human CD5. The screening of a rat cDNA library at high stringency using these probes resulted in a 1.5-kb positive clone. The DNA sequence of this clone confirmed its CD5 nature, but the clone appeared to lack part of the 5' and part of the 3' end. These missing 5' and 3' ends were obtained by PCR on rat thymus RNA. By ligating these PCR products to the original 1.5-kb CDM8 clone, a full-length rat CD5 gene was constructed. The full-length clone showed high identity with mouse and human CD5; however, at the 5' site of the gene a region of 36 nucleotides is present which is not seen in either mouse or human CD5. We have evidence that this sequence is a normal constituent of the rat CD5 gene: first, it is in frame with the rest of the CD5 coding sequence; second, it does not contain a stop codon; and third, it is also present in the CD5 gene of other rat strains. We transfected the full-length CD5 construct in COS cells and demonstrated that indeed the CD5 protein is recognized by MRC OX19. Although we showed that CD5 mRNA is present in rat B cells, extensive flow cytometry analysis using MRC OX19 as antibody failed to detect B cells expressing significant levels of CD5 on their cell surface compared to other B cells in any tissue or cell suspension tested from a variety of rat strains. This is in contrast with the mouse where a distinct population of B cells (B-1a cells) can be found expressing more CD5 than the other B cells. Either B-1 cells are not present in rats or CD5 is not the right phenotypic marker for rat B-1 cells. It still remains to be investigated whether a population of B cells (B-1a cells signer) in rats.

ANSWER 30 OF 66 MEDLINE

ANSWER 30 OF 66 MEDLINE

ACCESSION NUMBER: DOCUMENT NUMBER: 94166767 94166767

MEDLINE PubMed ID: 7907166 TITLE:

Chromosome 2-specific DNA clones from flow-sorted chromosomes of tomato.

AUTHOR: Arumuganathan K; Martin G B; Telenius H; Tanksley S D;

CORPORATE SOURCE: Department of Plant Breeding, Cornell University, Ithaca,

NY 14853-1902.
MOLECULAR AND GENERAL GENETICS, (1994 Mar) 242 (5) 551-8. SOURCE:

Journal code: NGP; 0125036. ISSN: 0026-8925. GERMANY: Germany, Federal Republic of Journal; Article; (JOURNAL ARTICLE)

PUB. COUNTRY:

LANGUAGE .

English Priority Journals FILE SEGMENT: ENTRY MONTH:

199404

SEGMENT: Priority Journals
199404
1Y MONTH: 199404
1Y DATE: Entered STN: 19940412
Last Updated on STN: 19950206
Entered Medline: 19940407
We obtained DNA clones specific to tomato chromosome 2 from a small number of chromosomes collected by flow sorting. Suspensions of metaphase chromosomes were prepared from 3-month-old tomato cell cultures of Lycopersicon pennellii. Isolated chromosomes stained with chromomycin A3 and Hoechst 33258 were analyzed on an EPICS 753 flow cytometer using a UV laser to excite theochst fluorescence and a 458 nm laser to excite chromomycin A3 fluorescence. Chromosomes from well-resolved peaks on a bivariate flow karyotype were sorted directly onto membrane filters for spot-blot analysis. The filters were processed and hybridized with chromosome-specific repetitive DNA probes. In this way tomato chromosome 1 and chromosome 2 were assigned to peaks in the bivariate flow karyotypes. One thousand copies of the putative chromosome 2 were flow-sorted directly into microfuge tubes. DNA specific to chromosome 2 derectly into microfuge tubes. DNA specific to chromosome 2 was amplified by a polymerase chain reaction (PCR) technique using universal 22mer degenerate oligonucleotide primers (DOP) sequences. DOP-PCR yields a smear of fragments of various sizes from 250 to 1600 bp. Amplified products were cloned into the Bluescript plasmid vector. Approximately 11% of the clones contained sequences with highly repetitive elements, and 85% contained only low-copy-number sequences. Eleven clones containing low-copy-number sequences that detect restriction fragment

length polymorphisms were placed on the molecular linkage map of tomato. All showed linkage to chromosome  $2\,.$ 

ANSWER 31 OF 66 MEDITINE

94180407 MEDLINE 94180407 PubMed ID: 8133535 ACCESSION NUMBER: DOCUMENT NUMBER:

Identification of partial complementary DNA clones encoding a 59-kd protein with characteristics of a unique oncofetal

antigen

Antigen.
Rashid H U; Barsoum A L; Cao T M; Coggin J H Jr
Department of Microbiology and Immunology, University of
South Alabama, College of Medicine, Mobile. AUTHOR: CORPORATE SOURCE:

CONTRACT NUMBER: CA39698 (NCI)

JOURNAL OF THE NATIONAL CANCER INSTITUTE, (1994 Apr 6) 86 (7) 515-26.

Journal code: J9J; 7503089. ISSN: 0027-8874. United States PUB. COUNTRY:

ANGUAGE:

Journal Article; (JOURNAL ARTICLE) English Priority Journals GENBANK-U06662 FILE SEGMENT: OTHER SOURCE: ENTRY MONTH: 199404 Entered STN: 19940428 ENTRY DATE:

Last Updated on STN: 19970203 Entered Medline: 19940420

BACKGROUND: Oncofetal antigens (OPAs) are conserved tumor-associated autoantigens or transplantation antigens present on the surface of all major classes of rodent and human tumors and on midgestational fetal cells but not on normal meonatal or adult human and rodent tissues. A but not on normal neonatal or adult human and rodent tissues. A syngeneically derived monoclonal antibody, MAD-115, recognizes murine OFAs of 44 and 200 kd in molecular mass. PURPOSE: Our goal was to clone and characterize the complementary DNAs (cDNAs) that encode these murine OFAs. METHODS: Rabbit antiserum raised against purified 44-kd OFA glycoprotein was used to screen a mouse embryo CDNA-lambda phage expression library. Recombinant phage clones positive for the expression of OFAs were detected by immunohistochemical staining, then isolated and plague purified. The presence of an OFA-encoding sequence in the plaque purified. The presence of an OFA-encoding sequence in the recombinant phage was confirmed by specific reaction of the expressed protein with MAb-115. Recombinant fusion protein was purified from the extracts of corresponding lysogens. Rabbit antiserum against purified recombinant fusion protein was raised, and the capacity of this antiserum recombinant fusion protein was raised, and the capacity of this antise to detect the expression of OFA on rodent tumor and fetal cells was determined by flow cytometry. In addition, immunoreactivity of tumor bearer and hyperimmune murine sera to bacterially expressed recombinant OFA protein was evaluated by enzyme-linked immunosorbent assay. The OFA-expressing insert DNA from plaque-purified lambda clones was subcloned into phagemid vectors for sequencing analysis. RESULTS: Antiserum derived against the isolated sequencing analysis. RESULTS: Antiserum derived against the isolated recombinant mouse embryo polypeptide mimicked MAD-115 in its specific binding to all OFA-positive rodent tumor and fetal cell lines tested and likewise did not show reactivity to normal adult tissues. This antiserum specifically recognized the native 44- and 200-kd OFAs in extracts of murine lymphocytic lymphoma. Furthermore, sera of tumor-bearing mice or mice immunized with purified OFA or intact, irradiated OFA-positive lymphocytic lymphoma cells also reacted with the recombinant fusion protein. The characterization of the isolated clone included nucleotide sequence information followed by analysis of the deduced primary structure of the protein. CONCLUSIONS: These data suggest that the isolated cDNA clones encode a distinct gene product which is widely expressed on the surface of tumor and fetal cells and represents the first characterized sequence of a true OFA. IMPLICATIONS: The availability of this cDNA, encoding a protein expressed only on tumor and fetal cells, provides a direct means to assess biological characteristics of malignant tissue which can be assayed by biochemical, histochemical, and molecular methods. which can be assayed by biochemical, histochemical, and molecular methods.

ANSWER 32 OF 66 MEDLINE DUPLICATE 9

94359457 MEDLINE 94359457 PubMed ID: 8078460 DOCUMENT NUMBER:

TITLE: Genome analysis of the moss Physcomitrella patens (Hedw.)

AUTHOR:

Reski R; Faust M; Wang X H; Wehe M; Abel W O Institut fur Allgemeine Botanik, Hamburg, Germany. MOLECULAR AND GENERAL GENETICS, (1994 Aug 15) 244 (4) CORPORATE SOURCE:

SOURCE:

Journal code: NGP; 0125036. ISSN: 0026-8925. GERMANY: Germany, Federal Republic of Journal; Article; (JOURNAL ARTICLE)

PUB. COUNTRY:

LANCHAGE .

FILE SEGMENT:

English
Priority Journals
GENBANK-X76632; GENBANK-X76634 OTHER SOURCE:

Entered STN: 19941013 ENTRY DATE:

Last Updated on STN: 19980206 Entered Medline: 19941006

Entered Medline: 19941006
A wild-type (WT) strain of the moss Physcomitrella patens (Hedw.) B.S.G., two mutants derived from it (PC22 and P24), and a somatic hybrid, PC22(+)P24, were analysed. Staining of metaphases revealed 54 +/- 2 chromosomes in the somatic hybrid and 27 chromosomes in the wild type and the two mutants. Using flow cytometry (FCM), DNA contents were calculated to be 0.6 pg (WT, PC22), 1.2 pg (P24), and 1.6 pg (PC22(+)P24) per nucleus, respectively. Southern hybridization provided evidence for at least one family of highly repetitive DNA and, furthermore, revealed different amounts of repetitive DNA in the four genotypes. However, these sequences cannot account for the 100% increase furthermore, revealed different amounts of repetitive DNA in the four genotypes. However, these sequences cannot account for the 100% increase in the nuclear DNA amount in mutant P24, relative to wild type. In FCM analyses every moss genotype generated just one single peak of fluorescence, indicating an arrest in the cell cycle during the daytime. Thermal denaturation of wild-type DNA revealed a G+c content of 34.6% for total DNA and 38.6% for plastid DNA. A cDNA library of 1.2 x 10(6) independent clones was established, from which sequences homologous to cab and rbcS, respectively, were isolated. These genes show significant homologies to those of higher plants, and, likewise, comprise multigene families. No restriction fragment length polymorphisms could be detected between the four moss genotypes using these cDNA probes.

detected between the four moss genotypes using these cDNA probes.

## 08/876,276 Search Strategy/Results

L5 ANSWER 33 OF 66 ACCESSION NUMBER: MEDLINE

94305578 94305578 MEDLINE PubMed ID: 7518285 DOCUMENT NUMBER:

TITLE

Jayosofa Funded ID: /S10202 Chromosome sorting by flow cytometry. Production of DNA libraries and gene mapping. Fantes J A; Green D K; Sharkey A Human Genetics Unit, Western General Hospital, Edinburgh,

AUTHOR:

CORPORATE SOURCE: Scotland.

METHODS IN MOLECULAR BIOLOGY, (1994) 29 205-19. Journal code: BU3; 9214969. ISSN: 1064-3745. SOURCE:

PUB. COUNTRY: United States

Journal; Article; (JOURNAL ARTICLE)

LANGUAGE: English Priority Journals

FILE SEGMENT: ENTRY MONTH: 199408

ENTRY DATE:

Entered STN: 19940825 Last Updated on STN: 19960129 Entered Medline: 19940816

ANSWER 34 OF 66 MEDLINE DUPLICATE 10

ACCESSION NUMBER: DOCUMENT NUMBER:

TITLE:

6 MEDLINE DUPLICATE 10
94101880 MEDLINE
94101880 PubMed ID: 8275710
A mouse chromosome 11 library generated from
sorted chromosomes using linker-adapter polymerase chain reaction.
Miyashita K; Vooijs M A; Tucker J D; Lee D A; Gray J W;

AUTHOR:

Pallavicini M G
Department of Laboratory Medicine, University of California, San Francisco 94103. CORPORATE SOURCE:

CONTRACT NUMBER: 27703

27703 CYTOGENETICS AND CELL GENETICS, (1994) 66 (1) 54-7. Journal code: DXK; 0367735. ISSN: 0301-0171. SOURCE:

PUB. COUNTRY:

Journal; Article; (JOURNAL ARTICLE) English Priority Journals LANGHAGE . FILE SEGMENT: ENTRY MONTH: 199402 Entered STN: 19940218

Y DATE:

Entered STN: 19940218

Last Updated on STN: 19940218

Entered Medline: 19940204

We describe the generation of a mouse whole chromosome library
using sequence-independent polymerase chain reaction (PCR) to amplify
sequences contained in DNA extracted from flow sorted chromosomes. DNA in
sorted chromosomes from a human x mouse hybrid cell line was digested with
a frequent four-cutter restriction enzyme, Sau3AI, and the ends were
ligated to an adapter oligonucleotide. The ligated DNA fragments were
amplified using PCR primers homologous to the linker-adapter
oligonucleotide. PCR-generated products were characterized by gel
electrophoresis. The size of the amplified DNA ranged from 100 to more
than 1,000 bp with a relatively high proportion of products at
approximately 400 bp. Biotinylated PCR products used for FISH showed
specific hybridization to murine metaphases and no hybridization to human
lymphocyte and hamster metaphase cells. Banding analysis indicated that
the probes were specific for mouse Chromosome 11. We anticipate that
availability of painting probes for specific murine chromosomes will
facilitate cytogenetic studies in the mouse.

ANSWER 35 OF 66 MEDLINE

ACCESSION NUMBER: DOCUMENT NUMBER: 93352474 93352474 MEDLINE

PubMed ID: 8394326

Human aminoacylase-1. Cloning, sequence, and expression analysis of a chromosome 3p21 gene inactivated in small TITLE:

cell lung cancer.
Cook R M; Burke B J; Buchhagen D L; Minna J D; Miller Y E

AUTHOR: CORPORATE SOURCE: Division of Pulmonary Sciences and Critical Care Medicine, University of Colorado Health Sciences Center, Denver 80262.

HL-07085 (NHLBI) CONTRACT NUMBER:

P50 CA58187 (NCI) RO1 HL45745 (NHLBI)

SOURCE: JOURNAL OF BIOLOGICAL CHEMISTRY, (1993 Aug 15) 268 (23)

Journal code: HIV; 2985121R. ISSN: 0021-9258. United States

Journal; Article; (JOURNAL ARTICLE) LANGUAGE:

PUB. COUNTRY:

English Priority Journals GENBANK-L07548 199309 FILE SEGMENT: OTHER SOURCE: ENTRY MONTH:

ENTRY DATE:

ON SOURCE: GENBANK-L07548

Y MONTH: 199309

Entered STN: 19931001

Last Updated on STN: 19970203

Entered Medline: 19930915

Human aminoacylase-1 (N-acyl-L-amino-acid amidohydrolase, EC 3.5.1.14;

ACYl) is a homodimeric zinc-binding enzyme that catalyzes the hydrolysis of N alpha-acylated amino acids. ACYl has been assigned to chromosome 3p21.1, a region reduced to homozygosity in small cell lung cancer (SCLC), and has been reported to exhibit reduced or absent expression in SCLC cell lines and tumors. Two human cDNA libraries and one human genomic DNA 11brary were screened with a previously isolated partial ACYl cDNA to isolate a full-length transcript. Sequence analysis of clones from each of these libraries resulted in an ACYl cDNA of 1438 base pairs with an open reading frame of 1224-base pairs coding for a putative protein of 408 amino acids with a predicted molecular mass of 45,882 Da. Sequence analysis revealed no homologies to previously reported cDNA or protein sequences and establishes ACYl as the first member of a new family of zinc-binding enzymes to be so characterized. The subcellular location of ACYl has been established as cytosolic by flow cytometry

Southern and northern analyses of ACYl in SCLC cell lines failed to demonstrate any gross abnormalities of the ACYl structural gene or instances of absent or aberrantly sized mRNA, respectively. instances of absent or aberrantly sized mRNA, respectively.

```
DOCUMENT NUMBER:
```

AUTHOR:

CORPORATE SOURCE:

94068469 PubMed ID: 7504279
Generation and screening of an oligonucleotide-encoded synthetic peptide library.
Needels M C; Jones D G; Tate E H; Heinkel G L;
Kochersperger L M; Dower W J; Barrett R W; Gallop M A
Affymax Research Institute, Palo Alto, CA 94304.
PROCEEDINGS OF THE NATIONAL ACADEMY OF SCIENCES OF THE
UNITED STATES OF AMERICA, (1993 Nov 15) 90 (22) 10700-4.
Journal code: PV3; 7505876. ISSN: 0027-8424.
United States
Journal; Atticle: (JOURNAL ARTICLE)

PUB. COUNTRY:

Journal; Article; (JOURNAL ARTICLE)

English FILE SEGMENT:

Priority Journals ENTRY MONTH: 199312 Entered STN: 19940201 ENTRY DATE:

Last Updated on STN: 19960129 Entered Medline: 19931229

Last Updated on STN: 19960129

Entered Medline: 19931229

We have prepared a library of approximately 10(6) different peptide sequences on small, spherical (10-microns diameter) beads by the combinatorial chemical coupling of both L- and D-amino acid building blocks. To each bead is covalently attached many copies of a single peptide sequence and, additionally, copies of a unique single-stranded oligonucleotide that codes for that peptide sequence. The oligonucleotide tags are synthesized through a parallel combinatorial procedure that effectively records the process by which the encoded peptide sequence is assembled. The collection of beads was screened for binding to a fluorescently labeled anti-peptide antibody using a fluorescence—activated cell sorting instrument. Those beads to which the antibody bound tightly were isolated by fluorescence—activated sorting, and the oligonucleotide identifiers attached to individual sorted beads were amplified by the PCR. Sequences of the amplified DNNs were determined to reveal the identity of peptide sequences that bound to the antibody with high affinity. By combining the capacity for information storage in an oligonucleotide code with the tremendous level of amplification possible through the PCR, we have devised a means for specifying the identity of each member of a vast library of molecules synthesized from both natural and unnatural chemical building blocks. In addition, we have shown that the use of flow cytometry incrementation parmits facile isolation of individual beads that hear addition, we have shown that the use of flow cytometry instrumentation permits facile isolation of individual beads that bear high-affinity ligands for biological receptors.

#### ANSWER 37 OF 66 MEDLINE

ACCESSION NUMBER: DOCUMENT NUMBER: 94077713 94077713 MEDLINE

PubMed ID: 8255765

TITLE: Coincidence painting: a rapid method for cloning region specific DNA sequences.

Erratum in: Nucleic Acids Res 1994 Feb 25;22(4):700 Bailey D M; Carter N P; de Vos D; Leversha M A; Perryman M COMMENT:

AUTHOR:

T; Ferguson-Smith M A CORPORATE SOURCE:

1; rerguson-minth M property (1993) Nov 11) 21 (22) 5117-23. NUCLEIC ACIDS RESEARCH, (1993 Nov 11) 21 (22) 5117-23. Journal code: 08L; 0411011. ISSN: 0305-1048. ENGLAND: United Kingdom SOURCE:

PUB. COUNTRY:

Journal; Article; (JOURNAL ARTICLE) English LANGUAGE:

FILE SEGMENT: Priority Journals ENTRY MONTH:

199401 ENTRY DATE: Entered STN: 19940203

Last Updated on STN: 19950206 Entered Medline: 19940113

Entered Medline: 19940113
We have developed a novel coincidence cloning strategy, termed Coincidence Painting, which enables the rapid generation of large numbers of region specific sequences. Coincidence Painting utilises Degenerate Oligonucleotide Primed PCR (DDP-PCR) amplification of flow sorted derivative translocation chromosomes. The PCR products are hybridised in situ onto specific flow sorted chromosomes for coincident sequence selection. Eluted and reamplified material is then cloned using a novel insert end revelation and ligation technique. Cloned inserts range in size from 150-1300 bps of which approximately 54% appear to be single copy sequences. The cloning method permits the excision of vector free probe for library hybridisation screening and the small insert size facilitates analysis for the generation of sequence tagged sites (STSs). We have used such clones successfully for YAC screening by PCR and for cosmid screening by filter hybridisation. This new methodology should allow the rapid saturation with probes of regions defined by specific translocation breakpoints. translocation breakpoints.

## EMBASE COPYRIGHT 2002 ELSEVIER SCI. B.V. 3356822 EMBASE ANSWER 38 OF 66

ACCESSION NUMBER: DOCUMENT NUMBER: 93356822

1993356822

TITLE: A strategy for the characterization of minute chromosome

A Birategy for the characterization or minute chromosome rearrangements using multiple color fluorescence in situ hybridization with chromosome-specific DNA libraries and YAC clones.

Popp S.; Jauch A.; Schindler D.; Speicher M.R.; Lengauer C.; Donis-Keller H.; Riethman H.C.; Cremer T. Inst Humangenetik und Anthropologie, Universitat Heidelberg, Im Neuenheimer Feld 328,D-69120 Heidelberg, Germany

AUTHOR:

CORPORATE SOURCE:

Germany Human Genetics, (1993) 92/6 (527-532). ISSN: 0340-6717 CODEN: HUGEDQ Germany SOURCE:

DOCUMENT TYPE: Journal; Article FILE SEGMENT: 022 Human Genetics

LANGUAGE: English SUMMARY LANGUAGE: English

The identification of marker chromosomes in clinical and tumor The identification of marker chromosomes in clinical and tumor cytogenetics by chromosome banding analysis can create problems. In this study, we present a strategy to define minute chromosomal rearrangements by multicolor fluorescence in situ hybridization (FISH) with 'whole chromosome painting' probes derived from chromosome-specific DNA libraries and Alu-polymerase chain reaction (PCR) products of various region-specific yeast artificial chromosome (YAC) clones. To demonstrate the usefulness of this strategy for the characterization of chromosome rearrangements unidentifiable by banding techniques, an 8p+ marker chromosome with two extra bands present in the karyotype of a child with multiple anomalies, malformations, and severe mental retardation was investigated. A series of seven-color FISH experiments with sets of fluorochrome-labeled DNA library probes from flow-sorted chromosomes demonstrated that the additional segment on 8p+ was derived from chromosome 6. For a more detailed characterization of the marker chromosome, three-color PISH experiments with library probes specific to chromosomes 6 and 8 were performed in combination with newly established telomeric and subtelomeric YAC clones from 6q25, 6p23, and 8p23. These experiments demonstrated a trisomy 6pter.fwdarw.6p22 and a monosomy 8pter.fwdarw.8p23 in the patient. The present limitations for a broad application of this strategy and its possible improvements are discussed.

ANSWER 39 OF 66 MEDLINE

MEDLINE PubMed ID: 8126985 ACCESSION NUMBER: 94172908 DOCUMENT NUMBER: 94172908

Human gene mapping and chromosome sorting by laser technology.

AUTTHOR:

Shimizu N; Minoshima S; Kudoh J; Kawasaki K; Wang Y; Shimizu Y; Sakai K

Department of Molecular Biology, School of Medicine, Keio University, Tokyo, Japan. KEIO JOURNAL OF MEDICINE, (1993 Dec) 42 (4) 212-6. Journal code: KUJ; 0376354. ISSN: 0022-9717. CORPORATE SOURCE:

SOURCE:

PUB. COUNTRY: Japan

Journal; Article; (JOURNAL ARTICLE) English

LANGUAGE: FILE SEGMENT: Priority Journals

ENTRY MONTH: 199404 Entered STN: 19940420 ENTRY DATE:

Last Updated on STN: 19940420 Entered Medline: 19940408

More than 30 human genes have been mapped by a combination of the fluorescent in situ hybridization (FISH) and spot blot hybridization in this laboratory for the past five years. Furthermore, the Keio strategy has been established for the molecular analysis of the human genome using flow-sorted human chromosomes.

ANSWER 40 OF 66 EMBASE COPYRIGHT 2002 ELSEVIER SCI. B.V. SSION NUMBER: 93040892 EMBASE ACCESSION NUMBER:

DOCUMENT NUMBER: 1993040892

Refined examination of plant metaphase chromosome structure at different levels made feasible by new isolation methods. Schubert I.; Dolezel J.; Houben A.; Scherthan H.; Wanner G. Institute of Plant Genetics, Crop Plant Research, 0-4325 Gatersleben, Germany Chromosoma, (1993) 102/2 (96-101). CORPORATE SOURCE:

Chromosoma, (1993) 102/2 (96-1 ISSN: 0009-5915 CODEN: CHROAU SOURCE:

COUNTRY: DOCUMENT TYPE: Germany

Journal; Article 022 English FILE SEGMENT: Human Genetics

LANGUAGE: SUMMARY LANGUAGE:

UAGE: English
ARY LANGUAGE: English
Two methods for isolation of plant metaphase chromosomes are described.
The first, micromanipulation, allows the isolation of a number of individual chromosomes, which may be used as templates for the generation of chromosome specific DNA libraries and for physical sequence mapping by the polymerase chain reaction (PCR). The second provides, from synchronized meristems, pure chromosome suspensions suitable for flow cytometric analysis and chromosome sorting. Restriction endonuclease banding, immunostaining of chromosomal antigens, as well as fluorescence in situ hybridization at high signal to noise ratio were successfully performed on the isolated chromosomes. Chromosomes obtained by both protocols were suitable for scanning electron microscopy, obtained by both protocols were suitable for scanning electron microscopy, the methods should also prove useful for refined analyses of the karyotypes of other plant species.

6 EMBASE COPYRIGHT 2002 ELSEVIER SCI. B.V. 93047361 EMBASE 1993047361 ANSWER 41 OF 66

ACCESSION NUMBER: DOCUMENT NUMBER:

TITLE:

1993047361
Identification of a tumor marker chromosome by flow sorting, DNA amplification in vitro, and in situ hybridization of the amplified product.
Boschman G.A.; Buys C.H.C.M.; Van der Veen A.Y.; Rens W.; Osinga J.; Slater R.M.; Aten J.A.
Academisch Medisch Centrum F0-212, Meibergdreef 9,1105 AZ Amsterdam, Netherlands
Genes Chromosomes and Cancer, (1993) 6/1 (10-16).
ISSN. 1045-2257, CODRN. GCCARS AUTHOR:

CORPORATE SOURCE:

ISSN: 1045-2257 CODEN: GCCAES United States

DOCUMENT TYPE: Journal; Article 016 Cancer FILE SEGMENT:

Human Genetics 022

English English LANGUAGE:

A method combining flow sorting and molecular cytogenetic techniques for the identification of unknown marker chromosomes is described. In this study, the bladder tumor cell line J82 was used, which was known to carry a marker chromosome of the size of chromosome 7 in every cell. From the cytogenetic analysis of Q-banded metaphase cells, it was shown to be composed of .apprx.40% presumably the greater part of chromosome 20 and for the rest microscopically unidentifiable material. This marker chromosome was found using flow cytometric analysis to form an independent peak and hence was suitable for isolation using dual-parameter sorting after staining with Hoechst 33258 and chromomycin A3. Subsequently, the marker was isolated by dual-parameter sorting. DNA amplification of 300 isolated chromosomes by polymerase chain reaction (PCR) using the Alu-primer Bk33 and the LINES-primer LH5 was carried out. After purification of the amplified product, a yield of 5 .mu.m of DNA was obtained. The DNA was labelled using Bio-II-dUTP and applied to human lymphocyte metaphase cells in a suppressive in situ hybridization procedure. Fluorescence was visible over chromosome 20 and over

the distal one-half of 6p. Together the fluorescent regions accounted for only .apprx.60% of the marker length, indicating a possible duplication of chromosome 20 material. This was confirmed by applying bicolor in situ hybridization using chromosome 6- and 20-specific DNA libraries to metaphase cells of the J82 cells.

L5 ANSWER 42 OF 66 ACCESSION NUMBER: MEDLINE

94001757 94001757 MEDLINE

DOCUMENT NUMBER:

94001757 PubMed ID: 7691162 Chromosomal radiosensitivity at intrachromosomal telomeric

sites.

Alvarez L; Evans J W; Wilks R; Lucas J N; Brown J M; Giaccia A J

CORPORATE SOURCE: Stanford University School of Medicine, Cancer Biology Research Laboratory, CA 94305-5468.

CA 55801 (NCI) CA 58838 (NCI) CONTRACT NUMBER:

GENES, CHROMOSOMES AND CANCER, (1993 Sep) 8 (1) 8-14. Journal code: AYV; 9007329. ISSN: 1045-2257. United States SOURCE:

PUB. COUNTRY:

Journal; Article; (JOURNAL ARTICLE)

LANGUAGE: FILE SEGMENT:

English

ENTRY MONTH:

ENTRY DATE:

Priority Journals 199311 Entered STN: 19940117 Last Updated on STN: 19960129 Entered Medline: 19931119

Entered Medline: 19931119

Telomeric DNA is composed of highly conserved sequences which are present at the termini of chromosomes as well as at intrachromosomal locations. Here, we studied a Chinese hamster ovary (CHO) cell line, BL-10, with highly stable amplified telomeric DNA at the termini as well as at intrachromosomal locations. We show that intrachromosomal or interstitial telomeric sites in this cell line and in another CHO cell line, HA-I, are radiosensitive in that they are more prone to breakage than would be expected based on the percentage of the genome composed of telomeric sequences. The frequency of breakage at interstitial telomeric sites is 4.3 to 8.3 times higher than that in the CHO genome overall. These conclusions are reached by both conventional cytogenetic analysis of two CHO cell lines which have the same survival rates after exposure to ionizing radiation, and by use of double fluorescence in situ hybridization (FISH) with a pan-telomere-specific probe and a CHO chromosome-specific library in the same metaphase cells after irradiation. irradiation

ANSWER 43 OF 66 MEDLINE

ACCESSION NUMBER: 92302264 MEDLINE

DOCUMENT NUMBER: 92302264 PubMed ID: 1376921

Random PCR mutagenesis screening of secreted proteins by Random PCR mutagenesis screening of secreted proteins by direct expression in mammalian cells.
Rice G C; Goeddel D V; Cachianes G; Woronicz J; Chen E Y; Williams S R; Leung D W
Department of Cell Biology, Genentech, Inc., South San Francisco, CA 94080.
PROCEEDINGS OF THE NATIONAL ACADEMY OF SCIENCES OF THE UNITED STATES OF AMERICA, (1992 Jun 15) 89 (12) 5467-71.
Journal code: PV3; 7505876. ISSN: 0027-8424.

CORPORATE SOURCE:

SOURCE:

PUB. COUNTRY: United States

Journal; Article; (JOURNAL ARTICLE)

LANGUAGE: English Priority Journals

AUTHOR:

FILE SEGMENT: ENTRY MONTH: 199207 ENTRY DATE:

SEGMENT: Priority Journals
YMONTH: 199207
PY MONTH: 199207

Entered STN: 19920731

Last Updated on STN: 19920721

We have developed a general method for screening randomly mutagenized expression libraries in mammalian cells by using fluorescence—activated cell sorting (FACS). The cDNA sequence of a secreted protein is randomly mutagenized by PCR under conditions of reduced Taq polymerase fidelity. The mutated DNA is inserted into an expression vector encoding the membrane glycophospholipid anchor sequence of decay-accelerating factor (DAF) fused to the C terminus of the secreted protein. This results in expression of the protein on the cell surface in transiently transfected mammalian cells, which can then be screened by FACS. This method was used to isolate mutants in the kringle 1 (K1) domain of tissue plasminogen activator (t-PA) that would no longer be recognized by a specific monoclonal antibody (mAb387) that inhibits binding of t-PA to its clearance receptor. DNA sequence analysis of the mutants and localization of the mutated residues on a three-dimensional model of the K1 domain identified three key discontinuous amino acid residues that are essential for mAb387 binding. Mutants with changes in any of these three residues were found to have reduced binding to the t-PA receptor on human hepatoma HepG2 cells but to retain full col lysis activity.

ANSWER 44 OF 66 EMBASE COPPRIGHT 2002 ELSPUER SCL B V

```
ANSWER 44 OF 66 EMBASE COPYRIGHT 2002 ELSEVIER SCI. B.V. SSION NUMBER: 92292171 EMBASE
```

ACCESSION NUMBER:

DOCUMENT NUMBER: 1992292171

Structural chromosome 1 aberrations in transitional cell

Structural chromosome 1 aberrations in transitional cell carcinoma of the bladder: Interphase cytogenetics combining a centromeric, telomeric, and library DNA probe.
Poddighe P.J.; Ramaekers F.C.S.; Smeets A.W.G.B.; Vooijs G.P.; Hopman A.H.N.
Molecular Cell Biol./Genetics Dept., University of Limburg, P.O. Box 616,6200 MD Maastricht, Netherlands
Cancer Research, (1992) 52/18 (4929-4934).
ISSN: 0008-5472 CODEN: CNREAB

AUTHOR:

CORPORATE SOURCE:

SOURCE:

United States Journal; Article 016 Cancer COUNTRY: DOCUMENT TYPE: FILE SEGMENT:

028

Human Genetics 022

Urology and Nephrology

English English LANGUAGE: SUMMARY LANGUAGE:

RRY LANGUAGE: ENGLISH
Fluorescence in situ hybridization (FISH) was used to study
numerical and structural chromosome 1 aberrations in interphase nuclei of

transitional cell carcinomas (TCCs) of the urinary bladder. One of the characteristic numerical aberrations, as detected previously in low-grade noninvasive TCCs, included trisomy for chromosome 1 (A. H. N. Hopman et al., Cancer Res., 51: 644-651, 1991). We examined in more detail 22 cases with a centromeric (1q12) and a telomeric associated (1p36) DNA probe and with a library DNA probe from sorted human chromosome 1 in single- and double-target FISH procedures. All flow cytometrically determined DNA diploid TCCs (13 cases), which showed three spots for 1q12 (6 cases), had two spots for 1p36. Since the library DNA probe showed three separate domains in the nuclei of these cases, the additional copy for 1q12 could be explained as an extra chromosome 1p-, containing the 1q12 target. In the flow cytometrically determined DNA tetraploid/aneuploid tumors, the results were more complex. In 6 of 9 cases, we observed an overrepresentation of 1q12 as compared to 1p36, also suggesting the presence of extra copies of 1p- chromosomes. The results of the present study demonstrate the utility of the FISH method to assess structural chromosome aberrations in interphase nuclei of solid tumors. transitional cell carcinomas (TCCs) of the urinary bladder. One of the

MEDLINE

MEDLINE

PubMed ID: 1565644

Cloning and expression of the tumor-associated antigen L6. Marken J S; Schieven G L; Hellstrom I; Hellstrom K E;

AUTHOR:

CORPORATE SOURCE:

Bristol-Myers Squibb Pharmaceutical Research Institute, Seattle, WA 98121. PROCEEDINGS OF THE NATIONAL ACADEMY OF SCIENCES OF THE UNITED STATES OF AMERICA, (1992 Apr 15) 89 (8) 3503-7. Journal code: PV3; 7505876. ISSN: 0027-8424. SOURCE:

PUB. COUNTRY: United States

Journal; Article; (JOURNAL ARTICLE)

LANGUAGE: English

FILE SEGMENT: OTHER SOURCE: ENTRY MONTH: Priority Journals GENBANK-M90657 199205

Entered STN: 19920607 Last Updated on STN: 19950206 Entered Medline: 19920515 ENTRY DATE:

The L6 cell surface antigen, which is highly expressed on lung, breast, colon, and ovarian carcinomas, has attracted attention as a therapeutic target for murine monoclonal antibodies and their humanized counterparts. target for murine monoclonal antibodies and their humanized counterparts. Its molecular nature has, however, remained elusive. Here we describe the expression cloning of a cDNA encoding the L6 antigen. COS cells transfected with this cDNA direct the expression of an approximately 24-kDa surface protein that reacts with the two anti-L6 monoclonal antibodies available. The predicted L6 peptide sequence is 202 amino acids long and contains three predicted NN2-terminal hydrophobic transmembrane regions, which are followed by a hydrophilic region containing two potential N-linked glycosylation sites and a COOH-terminal hydrophobic transmembrane region. The L6 antigen is related to a number of cell surface proteins with similar predicted membrane topology that have been implicated in cell growth. Two other members of this family of proteins, CD63 (MR491) and CO-029, are also highly expressed on tumor cells. The present findings should make it possible to further study the role of the L6-defined antigen in normal and neoplastic cells and to construct animal models for development of improved agents for active and passive cancer

models for development of improved agents for active and passive cancer

immunotherapy.

L5 ANSWER 46 OF 66 ACCESSION NUMBER: 6 MEDLINE 92242909

MEDLINE DOCUMENT NUMBER:

92242909 MEDLINE
92242909 PubMed ID: 1573270
A member of the tetra spans transmembrane protein superfamily is recognized by a monoclonal antibody raised against an HLA class I-deficient, lymphokine-activated killer-susceptible, B lymphocyte line. Cloning and preliminary functional studies.
Gil M L; Vita N; Lebel-Binay S; Miloux B; Chalon P; Kaghad M; Marchiol-Fournigault C; Conjeaud H; Caput D; Ferrara P;

AUTHOR:

CORPORATE SOURCE:

UAl156 Centre National de la Recherche Scientifique, Institut Gustave Roussy, Villejuif, France. JOURNAL OF IMMUNOLOGY, (1992 May 1) 148 (9) 2826-33. JOURNAL code: IFB; 2985117R. ISSN: 0022-1767. United States SOURCE:

PUB. COUNTRY:

Journal; Article; (JOURNAL ARTICLE)

LANGUAGE

FILE SEGMENT: Abridged Index Medicus Journals; Priority Journals ENTRY MONTH: 199206

ENTRY DATE:

Entered STN: 19920619

NONTH: 199206
RY DATE: Entered STN: 19920619
Last Updated on STN: 19980206 '
Entered Medline: 19920602

The IA4 mAb was identified among a series of antibodies raised in BALB/c mice after immunization against a HLA class I-deficient, lymphokine-activated killer (LAK)-susceptible EBV-B lymphocyte line. The IA4 antibody was selected because of its high expression, in the range of 10(5) to 25 x 10(5) sites/cell, on several B lymphocyte lines (EBV-transformed or Burkitt) and monocytic lines such as HL60 and U937, and because its expression was correlated with both target susceptibility to LAK lysis and reduced expression of HLA class I surface Ag on two pairs of EBV-b-transformed cell lines (721/721.134 and MM/10F2). Despite the strategy followed to raise the mAb and the correlation mentioned above, no direct role of the IA4 molecules in LAK susceptibility has been established, since the IA4 molecule is poorly expressed on the sensitive targets Daudi and K562; moreover, the IA4 antibody did not affect reproducibly the in vitro killing of positive target cells by LAK effectors. The IA4 antibody was poorly immunoprecipitating and the surface molecule recognized was identified by gene cloning following an expression strategy using a U937 cDNA library transfected in COS cells, and a screening strategy based on membrane expression of IA4 molecule. The IA4 cDNA is virtually identical to "R2." a mRNA species previously identified in activated human T cells by subtractive hybridization. The IA4 cDNA contains an open reading frame coding for a protein 267 amino acids long with four potential transmembrane domains and one large external hydrophilic domain of about 110 amino acids, possibly glycosylated. The

encoded protein belongs to a family of surface molecules, the tetra spans transmembrane protein superfamily, all displaying the four transmembrane domains, expressed on various cell types including lymphocytes (CD9, CD37, CD53, TAPA-1), melanoma cells (MFa91), and intestinal cells (CD0-C29). These molecules have been reported to be involved in cell activation and cell death. Surprisingly, the Schistosoma mansoni Ag Sm23 displays significant homologies with this family. The IA4 molecule is a widely distributed surface marker expressed on circulating lymphocytes and monocytes, newborn thymocytes, and the cell lines mentioned above. The IA4 molecule expression is up-regulated upon cell activation. Weakly expressed on resting peripheral T and B lymphocytes and large granular lymphocytes (NK), its expression roughly doubles after activation by PHA, staphylococcus aureus Cowan I, and IL-2, respectively.(ABSTRACT TRUNCATED AT 400 WORDS)

L5 ANSWER 47 OF 66 ACCESSION NUMBER:

6 MEDLINE 93078307 93078307 Pt MEDLINE DOCUMENT NUMBER: PubMed ID: 1447820

TITLE: Flow karyotyping and chromosome sorting. AUTHOR: Hashimoto K

CORPORATE SOURCE: Division of Genetic Resources, National Institute of

Health. SOURCE:

NIEPON RINSHO. JAPANESE JOURNAL OF CLINICAL MEDICINE, (1992 Oct) 50 (10) 2484-8. Ref: 9
Journal code: KIM; 0420546. ISSN: 0047-1852.

PUB. COUNTRY: Japan

Journal; Article; (JOURNAL ARTICLE)
General Review; (REVIEW)
(REVIEW LITERATURE)

LANGUAGE: FILE SEGMENT:

Japanese Priority Journals ENTRY MONTH: 199212

ENTRY DATE

AB

program was made to count pulses from the gated area only. As application of chromosome fractionation, flow cytometry has been used for karyotype analysis, gene mapping, and chromosomal DNA library construction.

DUPLICATE 12

L5 ANSWER 48 OF 66 MEDLINE
ACCESSION NUMBER: 92347875 MEDLINE
POCIMENT NUMBER: 92347875 PubMed ID: 1639403

Molecular mapping of mouse chromosomes 4 and 6: use of a flow-sorted Robertsonian chromosome. AUTHOR:

Bahary N; Pachter J E; Felman R; Leibel R L; Albright K; Cram S; Friedman J M

Cram S, Friedman J M
Howard Hughes Medical Institute, Rockefeller University,
New York, New York 10021.
DK26687 (NIDDK)
DK41096 (NIDDK)
HG00316 (NHGRI) CORPORATE SOURCE:

CONTRACT NUMBER:

GENOMICS, (1992 Jul) 13 (3) 761-9. Journal code: GEN; 8800135. ISSN: 0888-7543. SOURCE:

United States

PUB. COUNTRY:

Journal; Article; (JOURNAL ARTICLE) LANGUAGE:

English Priority Journals 199208 FILE SEGMENT: ENTRY MONTH: ENTRY DATE:

SEGMENT: Priority Journals
YMONTH: 199208
Entered STN: 1992091
Last Updated on STN: 19980206
Entered Medline: 19920828
The development of dense genetic maps of mammalian chromosomes is facilitated when chromosome-specific libraries are used as a source of genetic markers. To saturate the genetic maps of mouse chromosomes 4 and 6, we have made use of fluorescent-activated chromosome sorting . to purify a 4:6 Robertsonian chromosome from a cell line harboring the Rb(4:6)2Bnr translocation. After staining with chromomycin A3 and Hoechst 33528, this chromosome was separated from the other mouse chromosomes. DNA was isolated from the fraction containing the Robertsonian a hormosome and subcloned into the insertion vector lambda gt10, generating a library with 4.6 x 10(5) independent phage. A total of 19 single-copy sequences were used to type the progeny of a C57BL/6J x Mus spretus backcross that had previously been typed for loci on chromosomes 4 and 6. Approximately 70% of the clones in the library mapped to either chromosome 4 or 6 as assessed by genetic mapping and by use of a somatic cell hybrid panel. Simple sequence repeats have also been isolated from this library. Purther characterization of these microsatellites should accelerate efforts to map mouse chromosomes 4 and 6 using PCR. In addition, flow sorting of Robertsonian chromosomes suggests a general approach for making chromosome-specific libraries in mouse.

ANSWER 49 OF 66 MEDLINE DUPLICATE 13

MEDLINE DUPLICATE 13

ACCESSION NUMBER: 93052373 MEDLINE

DOCUMENT NUMBER:

TITLE:

93052373 MEDLINE
93052373 PubMed ID: 1427877
Mapping of 50 cosmid clones isolated from a flow-sorted human X chromosome library by fluorescence in situ hybridization.
Fan Y S; Sasi R; Lee C; Court D; Lin C C
Department of Laboratory Medicine and Pathology, University of Alberta, Edmonton, Canada.
GENOMICS, (1992 Oct) 14 (2) 542-5.
Journal code: GEN; 8800135. ISSN: 0888-7543.
United States CORPORATE SOURCE:

SOURCE:

PUB. COUNTRY:

United States
Journal; Article; (JOURNAL ARTICLE)

LANGUAGE: English

AUTHOR:

FILE SEGMENT: ENTRY MONTH: ENTRY DATE:

Priority Journals

199212 Entered STN: 19930122

Last Updated on STN: 19930122 Entered Medline: 19921201

Entered Medline: 19921201

Fifty cosmids have been mapped to metaphase chromosomes by

fluorescence in situ hybridization under conditions that suppress

signals from repetitive DNA sequences. The cosmid clones were isolated

from a flow-sorted human X chromosome library. Thirty-eight of

the clones were localized to chromosome X and 12 to autosomes such as

chromosomes 3, 7, 8, 14, and 17. Although most of the cosmids mapped to

the X chromosome appeared to be scattered along both the short and long

arms, 10 cosmids were localized to the centromeric region of the

chromosome. Southern blot analysis revealed that only two of these clones

hybridized to probe pXBR-1, which detects the DXZ1 locus. In addition, 4

out of 5 cosmids mapped on chromosome 8 also localized on the centromeric

region. While localization of X-specific cosmids will facilitate the

physical mapping of the human X chromosome, cosmids mapped to the

centromeric regions of chromosomes X and 8 should be especially useful for

studying the structure and organization of these regions.

ANSWER 50 OF 66 MEDLINE

ACCESSION NUMBER: DOCUMENT NUMBER: 93131264 93131264 MEDLINE

PubMed ID: 1483693

TITLE:

AUTHOR:

Complete characterization of a large marker chromosome by reverse and forward chromosome painting.
Blennow E; Telenius H; Larsson C; de Vos D; Bajalica S; Ponder B A; Nordenskjold M CORPORATE SOURCE:

Department of Clinical Genetics, Karolinska Institute, Stockholm, Sweden. HUMAN GENETICS, (1992 Dec) 90 (4) 371-4. Journal code: GED; 7613873. ISSN: 0340-6717. GERMANY: Germany, Federal Republic of Journal; Article; (JOURNAL ARTICLE) SOURCE: PUB. COUNTRY:

English LANGUAGE :

FILE SEGMENT:

Priority Journals

ENTRY MONTH: ENTRY DATE:

199302 Entered STN: 19930226 Last Updated on STN: 19930226 Entered Medline: 19930218

Entered Medline: 19930218

Marker chromosome are small supernumerary chromosomes that are sometimes associated with developmental abnormalities. Hence, the genes involved in such cases provide an interesting approach to understanding developmental abnormalities in man. As a first step towards isolating such sequences, marker chromosomes need complete characterization. By combining chromosome isolation by flow sorting and the "degenerate oligomucleotide primed - polymerase chain reaction", we have constructed a DNA library specific for a marker chromosome found in a child with severe developmental abnormalities. We used fluorescent in situ hybridization of the library onto normal metaphase spreads ("reverse chromosome painting") and were thus able to determine that the marker consists of the centromeric part of chromosome 7, the telomeric region of the long arm of chromosome 5 and the telomeric region of the short arm of the X-chromosome. Subsequently, we hybridized normal chromosome-specific libraries of the relevant chromosomes onto metaphases containing the marker chromosome "forward chromosome painting") and could in this manner establish the precise location of the different chromosome regions on the marker chromosome itself. This is a general approach suitable for outlining marker chromosomes in detail, and will aid the identification of the genes involved. identification of the genes involved.

ANSWER 51 OF 66 MEDLINE

ACCESSION NUMBER -92175525 92175525 MEDLINE

DOCUMENT NUMBER: PubMed ID: 1311700

TITLE:

Particles of a clone which induces expression of the gene encoding the human tumor necrosis factor receptor.

Garret M; McHendry-Rinde B; Spickofsky N; Margolskee R F Department of Neurosciences, Roche Research Center, Nutley, NJ 07110. AUTHOR: CORPORATE SOURCE:

SOURCE:

GENE, (1992 Feb 15) 111 (2) 215-22. Journal code: FOP; 7706761. ISSN: 0378-1119.

PUB. COUNTRY: Netherlands Journal; Article; (JOURNAL ARTICLE)

LANGUAGE: English

FILE SEGMENT:

Priority Journals
GENBANK-M64367; GENBANK-M64425; GENBANK-M64426;
GENBANK-M64427; GENBANK-M85145; GENBANK-S39996;
GENBANK-S40001; GENBANK-S40008; GENBANK-S40009; OTHER SOURCE

GENBANK-S40010

ENTRY MONTH:

199204 Entered STN: 19920424

Last Updated on STN: 19920424 Entered Medline: 19920408

Last Updated on STN: 19920424
Entered Medline: 19920408
Tumor necrosis factor (TNF) is a cytokine with pleiotropic effects upon cell growth, inflammation and immunologic responsiveness. High-affinity TNF receptors (TNFRs) of 55 and 75 kDa are found in many cell types. Using an Epstein-Barr virus (EBV)-based mammalian expression library, we have isolated a clone from human lymphoblastoid transfectants that induces overexpression of the TNFR-encoding gene (TNFR). Transfectants overproducing the TNFR were isolated by multiple rounds of sorting on a fluorescence-activated cell sorter using fluorescent TNF ligand binding as the selection procedure. Among the sorted transfectants were cells producing approx. 150,000 receptors per cell (Kd of approx. 1 nM). These cells have multiple copies of the TNFR gene present as extrachromosomal plasmids. These cells also overproduced the mRNA for TNFR. Low-Mr EBV episomes were isolated from these overproducing cells and used to transform Escherichia coli. One of the colonies isolated contained a plasmid encoding a portion of the noncoding region of the TNFR gene. Transfection of human lymphoblastoid cells with this DNA gave rise to high-level production of TNFR. Fluorescent TNF bound to these transfectants is fully and specifically displaced by an excess of TNF. The rescued clone contains approx. 10 kb of human genomic DNA including the 3'-untranslated region of TNFR and several Alu sequences; apparently during the selection procedure in human cells, recombination occurred to

rescue a portion of the TNFR gene. Transient transfection was used to narrow down the region responsible for TNFR induction to  $5.2~\rm kb$ . The mechanism by which this clone induces TNFR expression has not been determined.

ANSWER 52 OF 66 MEDLINE

92245414 92245414 ACCESSION NUMBER:

MEDLINE PubMed ID: 1315456 DOCUMENT NUMBER:

TITLE:

AUTHOR:

CORPORATE SOURCE:

CONTRACT NUMBER:

92245414 PubMed ID: 1315456
Cloning muscle isoforms of neural cell adhesion molecule using an episomal shuttle vector:
Pan L C; Margolskee R F; Blau H M
Pharmacology Department, Stanford University School of Medicine, California 94305.
HD18179 (NICHD)
HD20203 (NICHD)
SOMATIC CELL AND MOLECULAR GENETICS, (1992 Mar) 18 (2) SOURCE:

Journal code: UY2; 8403568. ISSN: 0740-7750.

United States PUB. COUNTRY: Journal; Article; (JOURNAL ARTICLE)

English Priority Journals LANGUAGE .

FILE SEGMENT: ENTRY MONTH: 199206 ENTRY DATE:

SEGENT: Priority Journals
Y MONTH: 199206
Y DATE: Entered STN: 19920619
Last Updated on STN: 19920619
Entered Medline: 19920602
Three distinct transcripts encoding two phosphatidylinositol (PI) linked isoforms of the neural cell adhesion molecule (NCAM) are induced during the differentiation of C2C12 myoblasts into myotubes. Corresponding NCAM clones were isolated from a mouse muscle cDNA library made in an Epstein-Barr virus shuttle vector that replicates extrachromosomally in human cells. Following transfection with the library, human cells expressing mouse NCAM were enriched using the fluorescence-activated cell sorter. Episomal NCAM clones recovered from sorted cells contain an 18-bp insert between exons 12 and 13. Two other NCAM cDNAs encode identical polypeptides containing a 108-bp insert homologous to the complete MSD1 domain, but differ in their 3' untranslated regions. Induction of MSD1-containing transcripts in advance of myotube formation suggests that muscle-specific NCAMs contribute to myogenesis from the earliest stages of differentiation. Moreover, our studies demonstrate the feasibility of cloning tissue-specific molecules by transfection and expression of cDNA libraries in episomal vectors.

ANSWER 53 OF 66 EMBASE COPYRIGHT 2002 ELSEVIER SCI. B.V. SSION NUMBER: 89275866 EMBASE MENT NUMBER: 1989275866

ACCESSION NUMBER: DOCUMENT NUMBER:

TITLE:

AUTHOR:

1989275866
Human chromosome-specific DNA libraries: Construction and purity analysis.
Fuscoe J.C.; McNinch J.S.; Collins C.C.; Van Dilla M.A. Biomedical Sciences Division, Lawrence Livermore National Laboratory, Livermore, CA 94550, United States
Cytogenetics and Cell Genetics, (1989) 50/4 (211-215).
ISSN: 0301-0171 CODEN: CGCGER CORPORATE SOURCE: SOURCE:

Switzerland Journal COUNTRY:

DOCUMENT TYPE: FILE SEGMENT: 022

Human Genetics 029 Clinical Biochemistry

English

SUMMARY LANGUAGE: English

ARY LANGUAGE: English
We report the construction of eight human chromosome-specific DNA
libraries. Metaphase chromosomes were purified by flow-sorting, and the
extracted DNA was cleaved with HindIII before cloning into lambda.Charon
21A. There is now a complete digest HindIII library containing
greater than five chromosome equivalents for each human chromosome. These
are available to the scientific community through the American Type
Culture Collection in Rockville, MD. The amount of hamster DNA in
libraries in which the chromosome was sorted from human x hamster hybrid
cells was estimated by species-specific hybridization. It ranged from 5%
to 39%. The sorted chromosomes were examined by fluorescence in
situ hybridization with species-specific DNA, and the main source of the
hamster DNA contamination was found to be intact hamster chromosomes. In
addition, we examined a chromosome 21 library, LL2INSO2, for
clones that fail to grow on the rec+ host LE392. Less than 0.6% of the
recombinant phage exhibited the rec+-inhibited phenotype.

ANSWER 54 OF 66 MEDLINE

ACCESSION NUMBER: 88320490 MEDLINE DOCUMENT NUMBER:

88320490 MEDLINE
88320490 PubMed ID: 3413114
A highly conserved repetitive DNA sequence, (TTAGGG)n, present at the telomeres of human chromosomes.
Moyais R K; Buckingham J M; Cram L S; Dani M; Deaven L L; Jones M D; Meyne J; Ratliff R L; Wu J R
Genetics Group, LS-3, Los Alamos National Laboratory, University of California 87545.
RR01315 (NCPR) AUTHOR:

CORPORATE SOURCE:

RR01315 (NCRR) CONTRACT NUMBER:

PROCEEDINGS OF THE NATIONAL ACADEMY OF SCIENCES OF THE

UNITED STATES OF AMERICA, (1988 Sep) 85 (18) 6622-6.

Journal code: PV3; 7505876. ISSN: 0027-8424.

United States

PUB. COUNTRY:

Journal; Article; (JOURNAL ARTICLE) English

LANGUAGE: FILE SEGMENT:

Priority Journals GENBANK-J04078; GENBANK-J04078; GENBANK-M19946 OTHER SOURCE: ENTRY MONTH:

ENTRY DATE:

TITLE:

Entered STN: 19900308 Last Updated on STN: 19970203 Entered Medline: 19881013

Entered Medline: 19881013
A highly conserved repetitive DNA sequence, (TTAGGG)n, has been isolated from a human recombinant repetitive DNA library. Quantitative hybridization to chromosomes sorted by flow cytometry indicates that comparable amounts of this sequence are present on each human chromosome. Both fluorescent in situ hybridization and BAL-31 nuclease digestion experiments reveal major clusters of this sequence at the telomeres of all human chromosomes. The evolutionary

conservation of this DNA sequence, its terminal chromosomal location in a variety of higher eukaryotes (regardless of chromosome number or chromosome length), and its similarity to functional telomeres isolated from lower eukaryotes suggest that this sequence is a functional human relumence. telomere.

L5 ANSWER 55 OF 66 MEDLINE ACCESSION NUMBER: 87260984 DOCUMENT NUMBER: 87260984 Pu MEDLINE

87260984 PubMed ID: 2955415
Carcinoembryonic antigen family: expression in a mouse
L-cell transfectant and characterization of a partial cDNA

AUTHOR:

SOURCE:

L-Cell transfectant and characterization of a partial c in bacteriophage lambda gtil.

Kamarck M E; Elting J J; Hart J T; Goebel S J; Rae P M; Nothdurft M A; Nedwin J J; Barnett T R PROCEEDINGS OF THE NATIONAL ACADEMY OF SCIENCES OF THE UNITED STATES OF AMERICA, (1987 Aug) 84 (15) 5350-4.

Journal code: PV3; 7505876. ISSN: 0027-8424.

PUB. COUNTRY: United States

Journal; Article; (JOURNAL ARTICLE) English

LANGUAGE:

FILE SEGMENT: OTHER SOURCE: Priority Journals GENBANK-M17191 ENTRY MONTH: 198708

ENTRY DATE:

RESOURCE: GENBANK-M17191

YMONTH: 198708

Entered STN: 19900305

Entered Medline: 19870826

Genomic DNA and mRNA from the adenocarcinoma cell line LoVo were used to generate L-cell transfectants and a bacteriophage lambda gtll cDNA clone that express epitopes of carcinoembryonic antigen (CEA). Primary and secondary L-cell transfectants expressing CEA were selected with a fluorescence-activated cell sorter (FACS). These transfectants, including some clones that were selected for high-level CEA expression by multiple rounds of FACS sorting, express a surface protein of 150 kDa that reacts with all anti-CEA antibodies tested. In parallel, a cDNA library of LoVo poly(A)+ RNA was constructed in lambda gtll and fusion proteins were screened with polyclonal antisera against CEA. One positive clone, lambda cLV7, was identified that hybridized specifically to transfectant DNA. The nucleic acid sequence of the cDNA insert (cLV7) contained two regions of extensive internal homology, with greater than 70% identity at the amino acid level. cLV7 hybridized to three mRNA species of LoVo cells and to a predominant mRNA of the CEA-expressing transfectants. Hybridization of cLV7 to restriction endonuclease-digested genomic DNA of colon carcinoma cells, normal human cells, and human-mouse somatic cell hybrids revealed the presence of multiple hybridizing bands, one of which was present in transfectant cells. These CEA-related sequences are not rearranged in tumors and, by somatic cell hybrid analysis, were mapped to human chromosome 19.

ANSWER 56 OF 66 MEDLINE DUPLICATE 15 AB

ANSWER 56 OF 66 MEDLINE

ACCESSION NUMBER: DOCUMENT NUMBER: 88054312 MEDLINE 88054312 PubMed ID: 3677921

TITLE.

88054312 PubMed ID: 3677921 Human chromosome-specific repetitive DNA sequences: novel markers for genetic analysis. Moyais R K; Albright K L; Bartholdi M F; Cram L S; Deaven L L; Hildebrand C E; Joste N E; Longmire J L; Meyne J; Schwarzacher-Robinson T AUTHOR:

DUPLICATE 15

CORPORATE SOURCE:

Schwarzacher-Robinson T Genetics Group, Los Alamos National Laboratory, University of California, NM 87545. RR01315 (NCRR) CHROMOSOMA, (1987) 95 (6) 375-86. Journal code: D7A; 2985138R. ISSN: 0009-5915. GERMANY, WEST: Germany, Federal Republic of Journal; Article; (JOURNAL ARTICLE) English

CONTRACT NUMBER:

SOURCE:

PUB. COUNTRY:

LANGUAGE:

English Priority Journals FILE SEGMENT:

OTHER SOURCE: ENTRY MONTH: GENBANK-M20262; GENBANK-M20263

ENTRY DATE:

Entered STN: 19900305

IN DATE: 19472

IF DATE: Entered STN: 19900305

Last Updated on STN: 19970203

Entered Medline: 19871231

Two recombinant DNA clones that are localized to single human chromosomes were isolated from a human repetitive DNA library. Clone pHuR 98, a variant satellite 3 sequence, specifically hybridizes to chromosome position 9qh. Clone pHuR 195, a variant satellite 2 sequence, specifically hybridizes to chromosome position 16qh. These locations were determined by fluorescent in situ hybridization to metaphase chromosomes, and confirmed by DNA hybridizations to human chromosomes sorted by flow cytometry. Pulsed field gel electrophoresis analysis indicated that both sequences exist in the genome as large DNA blocks. In situ hybridization to intact interphase nuclei showed a well-defined, localized organization for both DNA sequences. The ability to tag specific human autosomal chromosomes, both at metaphase and in interphase nuclei, allows novel molecular cytogenetic analyses in numerous basic research and clinical studies.

EMBASE COPYRIGHT 2002 ELSEVIER SCI. B.V. 6209738 EMBASE ANSWER 57 OF 66

ACCESSION NUMBER: 86209738

DOCUMENT NUMBER:

AUTHOR: CORPORATE SOURCE:

86209738 EMBASE
1986209738
Cloning of DNA libraries from mouse Y chromosomes purified by flow cytometry.
Baron B.; Metezeau P.; Hatat D.; et al.
Unite d'Immunogenetique Humaine (Inserm U.276), Institut Pasteur, 75724 Paris Cedex 15, France
Somatic Cell and Molecular Genetics, (1986) 12/3 (289-295).
CODEN: SCMCDN: SOURCE:

COUNTRY -United States DOCUMENT TYPE: Journal

FILE SEGMENT: 022 Human Genetics

English

To purify mouse Y chromosomes by flow cytometry, a male cell line containing the Robertsonian translocation Rb(9.19)163H has been established by SV40 transformation. Flow karyotypes obtained from these cells exhibit a well-isolated peak of fluorescence

corresponding to the single Y chromosome, clearly distinct from that of

chromosome 19. From this peak, 650,000 chromosomes were sorted, and two restriction fragment libraries were constructed from the DNA of the sorted chromosomes. The characterization of several Y-specific fragments has shown that the Y DNA was enriched at least 36-fold. Furthermore, given that there are likely homologies between the X and Y chromosomes, we can assume that this calculated value of the purification factor is an underestimation and that the Y DNA was more highly purified by flow sorting. ANSWER 58 OF 66 MEDLINE SSION NUMBER: 87132326 MEDLINE DUPLICATE 16 ACCESSION NUMBER: 87132326 PubMed ID: 3816174 SCAN: a program library for high resolution slit-scan analysis of chromosomes in flow DOCUMENT NUMBER: TITLE: cytometry.
Weier H U; Eisert W G
COMPUTER METHODS AND PROGRAMS IN BIOMEDICINE, (1986 Dec) 23 AUTHOR:

SOURCE:

(3) 269-76. Journal code: DOH; 8506513. ISSN: 0169-2607.

PUB. COUNTRY: Netherlands

Journal; Article; (JOURNAL ARTICLE) English

LANGUAGE: FILE SEGMENT:

Priority Journals 198703

ENTRY DATE:

Entered STN: 19900303

Last Updated on STN: 19900303 Entered Medline: 19870330

Entered Medline: 19870330

Computer programs for high resolution slit-scan chromosome analysis in flow cytometry are introduced. A modular program library, SCAN, contains programs for single and dual parameter data acquisition, correction of recorded profiles and computation of histograms of various parameters. Using a minicomputer system, data acquisition is programmed in assembler to realize high input rates and real-time histogram calculation. Software for the processing of recorded profiles has been written in FORTRAN and allows extensions or alterations for different objectives. A sample run recording bicolor fluorescence profiles from metaphase chromosomes demonstrates the main features of the software. main features of the software.

EMBASE COPYRIGHT 2002 ELSEVIER SCI. B.V. 5063328 EMBASE

L5 ANSWER 59 OF 66 EMBASE ACCESSION NUMBER: 85063328

DOCUMENT NUMBER: 1985063328

P98063328 Flow-cytometric analysis and sorting of human metaphase chromosomes. A new field of research.
Tranebjaerg L.; Vissing H.; Mikkelsen M.
Kromosomlaboratoriet, John F. Kennedy Instituttet, DK-2600 AUTHOR:

CORPORATE SOURCE:

Glostrup, Denmark Ugeskrift for Laeger, (1985) 147/4 (239-242). SOURCE: CODEN: UGLAAD

Denmark

COUNTRY: DOCUMENT TYPE: Journal

FILE SEGMENT: 022 Human Genetics LANGUAGE: Danish

SUMMARY LANGUAGE:

RRY LANGUAGE: English
Flow-cytometric characterisation of suspended metaphase chromosomes Flow-cytometric characterisation of suspended metaphase chromosomes obtained from various cultured human cells is feasible for karyotyping purposes. Certain advantages of flow-cytometry applied as a diagnostic tool in cytogenetics are presented: 1) Quick, reproducible analysis of a large number of cells (106-107). 2) The resulting flow-histogram is constant for each individual patient. On the other hand, some limitations of diagnostic flow cytometry exist:

1) Consideration of the patient (a large volume of blood is necessary to obtain a sufficient number of lymphocytes). 2) Pronounced inter-individual variation in the flow-histograms obtained from normal persons is the consequence of heterochromatic polymorphism for a number of chromosomes.

3) Precise statement of specific chromosomes to specific peaks of the fluorescence-histograms presumes visual (in fluorescence microscope) analysis of individual chromosome fractions.

Fluorescence-activated chromosome sorting (FACS) is a feasible method of isolating specific chromosomes in sufficient numbers for biochemical studies of DNA. By means of FACS is it possible: 1) to establish libraries of DNA-sequences from specific chromosomes (genelibrary) as already reported concerning chromosome X, Y, 21 and library) as already reported concerning chromosome X, Y, 21 and 22; 2) to map genes to individual chromosomes or specific parts of chromosomes; and 3) to isolate otherwise unidentified marker chromosomes from tumours in order to investigate whether DNA-sequence homology with identified chromosomes can be found.

ANSWER 60 OF 66 EMBASE COPYRIGHT 2002 ELSEVIER SCI. B.V. SSION NUMBER: 85170360 EMBASE

ACCESSION NUMBER:

DOCUMENT NUMBER: 1985170360

TITLE: Human chromosome analysis by flow

AUTHOR:

Human chromosome analysis by flow cytometry.
Young B.D.
Medical Oncology Laboratory, Imperial Cancer Research Fund,
St. Bartholomew's Hospital, London EC1, United Kingdom
Annals of the New York Academy of Sciences, (1985) VOL.
450/- (11-23).
CODEN: ANYAA
United States CORPORATE SOURCE:

SOURCE:

COUNTRY: Journal

DOCUMENT TYPE: FILE SEGMENT: Human Genetics 022

English LANGUAGE:

UAGE: English

There have been many attempts to fractionate metaphase chromosomes by centrifugation, countercurrent distribution, and 1g sedimentation. While such methods can separate chromosomes from the other cellular constituents, and have achieved enrichment of different size classes of chromosomes, the purification of single chromosomes was not possible. Only in recent years has it become possible to fractionate chromosomes with a high degree of resolution. The application of the fluorescence activated cell sorter has allowed the purification of small but useful quantities of individual chromosomes, and therefore has become the method of choice for tackling a number of biological questions. With recombinant DNA techniques, it is now also possible to generate a large number of DNA

probes for a single flow-sorted chromosome and this will prove extremely useful in the linkage analysis of certain genetic disorders.

ANSWER 61 OF 66 MEDLINE

ACCESSION NUMBER: 84181960 84181960 MEDLINE

PubMed ID: 6232119 DOCUMENT NUMBER:

84181960 PubMed ID. 6232119
Development and use of metaphase chromosome flow-sorting methodology to obtain recombinant phage libraries enriched for parts of the human X chromosome.
Lalande M; Kunkel L M; Flint A; Latt S A
GM21121/33579 (NIGMS)
HD04807/18658 (NICHD)
CYTOMETRY, (1984 Mar) 5 (2) 101-7.
Journal code: D92; 8102328. ISSN: 0196-4763.
United States '
Journal: Article: (JOURNAL ARTICLE)

AUTHOR:

CONTRACT NUMBER:

SOURCE:

PUB. COUNTRY:

Journal; Article; (JOURNAL ARTICLE)

LANGUAGE: English FILE SEGMENT: ENTRY MONTH: ENTRY DATE: Priority Journals 198406

Entered STN: 19900319

Last Updated on STN: 19970203 Entered Medline: 19840619

Last Updated on STN: 19970203

Entered Medline: 19840619

Metaphase chromosomes isolated from human lymphoblastoid cell lines containing structurally abnormal X chromosomes have been stained with the bisbenzimidazole dye Hoechst 33258 and analyzed on a PACS II flow system equipped with a 5-W all-lines argon ion laser. The chromosomal fluorescence has been highly resolved at flow rates of 1,000-3,000 chromosomes per second. With the goal of obtaining recombinant DNA libraries from parts of the human X chromosome, fluorescence populations enriched for a dicentric X (Xpter-greater than Xq24::Xq24-greater than Xpter) chromosome and an isochromosome of the long arm of the X [i(Xq)] have been identified. The dicentric X chromosome has been resolved as a discrete peak in the fluorescence flow histogram. In contrast, the fluorescence intensity of the isochromosome is indistinguishable from that of chromosomes 3 and 4. Recombinant DNA libraries from the flow-sorted chromosomes have been constructed in the lambda phage, Charon 21A, and consist of 1.6 X 10(5) and 0.7 X 10(5) plaque-forming units in the case of the dicentric X and the isochromosome, respectively. Ninety percent of the phage in both recombinant libraries contain inserts which hybridize to highly repetitive human DNA sequences. The recombinant phage library from the flow-sorted dicentric X chromosome, which could be assigned to a discrete fluorescence peak, has been further characterized and shows at least a tenfold enrichment for X chromosome-specific DNA sequences as determined by Southern blot hybridization of cloned fragments.

L5 ANSWER 62 OF 66 ACCESSION NUMBER:

DOCUMENT NUMBER: TITLE:

AUTHOR:

CONTRACT NUMBER:

SOURCE:

86 MEDLINE
83247418 MEDLINE
83247418 PubMed ID: 6575396
Isolation of amplified DNA sequences from IMR-32 human neuroblastoma cells: facilitation by fluorescence
-activated flow sorting of metaphase chromosomes.
Kanda N; Schreck R; Alt F; Bruns G; Baltimore D; Latt S CA-14501 (NCI)
CA-26717 (NCI)
HD-04807 (NICHD)
PROCEEDINGS OF THE NATIONAL ACADEMY OF SCIENCES OF THE UNITED STATES OF AMERICA, (1983 Jul) 80 (13) 4069-73.
Journal code: PV3; 7505876. ISSN: 0027-8424.
United States

PUB. COUNTRY: United States

Journal; Article; (JOURNAL ARTICLE)

LANGUAGE:

English Priority Journals FILE SEGMENT: Entered STN: 19900319 ENTRY DATE:

Last Updated on STN: 19970203 Entered Medline: 19830811

Entered Medline: 19830811

Human neuroblastoma IMR-32 cells have large homogeneously staining regions (HSRs), primarily in the short arms of chromosome 1. We have constructed a recombinant phage library that is enriched for DNA present in the HSR of this chromosome by using fluorescence-activated flow sorting for initial chromosome purification. Eleven distinct cloned DNA segments were identified that showed significantly greater hybridization to IMR-32 genomic DNA, detected by Southern blotting, than to normal human genomic DNA. These sequences have also been localized to the HSR of chromosome 1 by in situ hybridization. Based on an approximate 50-fold sequence amplification for each cloned segment and a total HSR size of 150,000 kilobases, the amplified unit in the HSR is estimated to be 3,000 kilobases. Sequences homologous to all cloned HSR DNA segments were mapped to human chromosome 2 by using human-mouse hybrid cells. Further work using in situ hybridization demonstrated that cloned HSR segments were localized in the short arm of chromosome 2 in both normal and IMR-32 cells may have involved transposition from chromosome 2 to chromosome I.

ANSWER 63 OF 66 MEDLINE

ACCESSION NUMBER: 84003819 84003819 MEDLINE

DOCUMENT NUMBER: TITLE:

SOURCE:

84003819 MEDLINE
84003819 PubMed ID: 6193931
Purifying human Y chromosomes by flow
cytometry and sorting.
Fantes J A; Green D K; Cooke H J
CYTOMETRY, (1983 Jul) 4 (1) 88-91.
Journal code: D92; 8102328. ISSN: 0196-4763.

PUB. COUNTRY: United States

Journal; Article; (JOURNAL ARTICLE)

LANGUAGE: English

FILE SEGMENT: Priority Journals ENTRY MONTH: ENTRY DATE:

AB

individuals. A suspension of chromosomes in a polyamine buffer was individuals. A suspension of chromosomes in a polyamine buffer was produced from each sample, stained with the fluorescent dye Hoechst 33258, and passed through a flow cytometer and sorter. Following analysis of the 17 fluorescence distributions, a single donor was found giving a separate peak corresponding to the Y chromosome. Seventy percent of the chromosomes sorted from this peak were identified as Y chromosomes. Batches of a million Y chromosomes were produced from each of several 40 ml donations of peripheral blood. These were assessed for the amount of Y DNA present and used to construct a DNA library.

6 MEDLINE 82222258 DUPLICATE 17

MEDLINE ACCESSION NUMBER:

DOCUMENT NUMBER:

82222258 PubMed ID: 6953442 Construction and characterization of genomic libraries from TITLE:

Construction and characterization of genomic libraries specific human chromosomes.

Krumlauf R; Jeanpierre M; Young B D

PROCEEDINGS OF THE NATIONAL ACADEMY OF SCIENCES OF THE UNITED STATES OF AMERICA, (1982 May) 79 (9) 2971-5.

Journal code: PV3; 7505876. ISSN: 0027-8424. AUTHOR: SOURCE:

PUB. COUNTRY: United States

Journal; Article; (JOURNAL ARTICLE)

LANGUAGE :

English Priority Journals FILE SEGMENT: ENTRY MONTH: 198208 ENTRY DATE: Entered STN: 19900317 Last Updated on STN: 19900317

Entered Medline: 19820814
Highly purified fractions of human chromosomes 21 and 22 were isolated from a suspension of metaphase chromosomes stained with ethidium bromide by using a fluorescence-activated cell sorter (FACS II). Two by using a fluorescence activated cell sorter (FACS II). Two recombinant DNA libraries, representing chromosomes 21 and 22, were constructed by complete digestion of DNA from these fractions with EcoRI and insertion into the vector lambda gtWES lambda B. Twenty clones selected at random from the chromosome 22 library hybridized to EcoRI-digested human DNA, and five of these clones hybridized to single bands identical in size to the phage inserts. These five single-copy sequences and a clone coding for an 85 RNA isolated by screening the chromosome 22 library for expressed sequences were characterized in detail. Hybridization of all six clones to a panel of sorted chromosomes and hybrid cell lines confirmed the assignment of the sequences to chromosome 22. The sequences were localized to regions of chromosome 22 by hybridization to translocated chromosomes sorted from a cell line having a balanced translocation (17;22) (pl3;q11) and to hybrid cell lines containing the various portions of another translocation t(X;22) (q13;q112). Five clones reside on the long arm of chromosome 22 between q112 and pter, while one clone and an 185 rRNA gene isolated from the chromosome-specific libraries by this method has the advantage of being direct and applicable to nearly all human chromosomes and will be important in molecular analysis of human genetic diseases.

ANSWER 65 OF 66 MEDLINE

ACCESSION NUMBER: DOCUMENT NUMBER: 82174330 82174330 MEDITINE

TITLE:

82174330 MEDLINE
82174330 PubMed ID: 6461845
Regional localization on the human X of DNA segments cloned from flow sorted chromosomes.
Kunkel L M; Tantravahi U; Eisenhard M; Latt S A GM21121 (NIGMS)
HD 40807 (NICHD)
NUCLEIC ACIDS RESEARCH, (1982 Mar 11) 10 (5) 1557-78.
JOURNAL code: 08L; 0411011. ISSN: 0305-1048.
ENGLAND: United Kingdom
Journal; Article; (JOURNAL ARTICLE)

AUTHOR:

important in molecular analysis of human genetic diseases.

CONTRACT NUMBER:

SOURCE:

PUB. COUNTRY:

LANGUAGE: English

FILE SEGMENT: Priority Journals 198206 Entered STN: 19900317 ENTRY DATE: Last Updated on STN: 19970203 Entered Medline: 19820614

Entered Medline: 19820614

Fluorescence activated sorting of chromosomes from 49,XXXXY
human lymphoblasts has been used to obtain DNA enriched for the human X.
This DNA was cloned in lambda phage Charon 21A to obtain a library
of approximately 60,000 pfu. Phage inserts free of human highly repeated
DNA sequences are localized to different regions of the human X by two
independent hybridization analyses. The first utilized comparative
hybridization to rodent-human hybrid cell DNA samples containing all or
known portions of the human X, while the second was based on hybridization
dosage to DNA samples from human cell lines differing in the number of X
chromosomes or X chromosome segments. Of five unique sequence inserts
tested, three were X chromosome specific and were localized to regions
Xpter leads to Xcen, Xql leads to Xq22 and Xq24 leads to Xq24 leads to Xq22. Xpter leads to Xcen, Xql leads to Xq22 and Xq24 leads to Xqter, respectively. The library presented here represents a highly enriched source of human X chromosome-specific DNA sequences.

ANSWER 66 OF 66 MEDLINE DUPLICATE 18 ACCESSION NUMBER: 82013644 MEDLINE

DOCUMENT NUMBER:

82013644

82013644

PubMed ID: 6456416

Cloning of a representative genomic library of the human X chromosome after sorting by flow TITLE:

cytometry.
Davies K E; Young B D; Elles R G; Hill M E; Williamson R
NATURE, (1981 Oct 1) 293 (5831) 374-6.
Journal code: NSC; 0410462. ISSN: 0028-0836. AUTHOR:

SOURCE:

PUB. COUNTRY:

ENGLAND: United Kingdom
Journal; Article; (JOURNAL ARTICLE)

LANGUAGE: English Priority Journals

FILE SEGMENT: ENTRY MONTH: 198111

ENTRY DATE:

## 08/876,276 Search Strategy/Results

separated using a fluorescence-activated cell sorter. The DNA was purified from the chromosomes, digested to completion with the restriction enzyme EcoRI and cloned into the phage lambda gtWES.lambda B. The X-derived nature of the recombinants was confirmed by hybridization to rodent/human cell line DNA containing only the human X chromosome. Such libraries will be particularly useful for the investigation of genetic diseases such as Duchenne muscular dystrophy, where the basic defect has not been elucidated, and of neoplasia, where several specific chromosomal anomalies, particularly for the leukaemias, have been identified.